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(54) Title: HIGH LEVEL EXPRESSION OF PROTEINS

(57) Abstract

The invention features a synthetic gene encoding a protein normally expressed in a mammalian cell wherein at least one non-preferred or less preferred codon in the natural gene encoding the protein has been replaced by a preferred codon encoding the same amino acid.

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HIGH LEVEL EXPRESSION OF PROTEINS

Field of the Invention

The invention concerns genes and methods for expressing eukaryotic and viral proteins at high levels in eukaryotic cells.

Background of the Invention

Expression of eukaryotic gene products in prokaryotes is sometimes limited by the presence of codons that are infrequently used in *E. coli*. Expression of such genes can be enhanced by systematic substitution of the endogenous codons with codons over represented in highly expressed prokaryotic genes (Robinson et al., Nucleic Acids Res. 12:6663, 1984). It is commonly supposed that rare codons cause pausing of the ribosome, which leads to a failure to complete the nascent polypeptide chain and a uncoupling of transcription and translation. Pausing of the ribosome is thought to lead to exposure of the 3' end of the mRNA to cellular ribonucleases.

Summary of the Invention

The invention features a synthetic gene encoding a protein normally expressed in a mammalian cell or other eukaryotic cell wherein at least one non-preferred or less preferred codon in the natural gene encoding the protein has been replaced by a preferred codon encoding the same amino acid.

Preferred codons are: Ala (gcc); Arg (cgc); Asn (aac); Asp (gac) Cys (tgc); Gln (cag); Gly (ggc); His (cac); Ile (atc); Leu (ctg); Lys (aag); Pro (ccc); Phe (ttc); Ser (agc); Thr (acc); Tyr (tac); and Val (gtg). Less preferred codons are: Gly (ggg); Ile (att); Leu (ctc); Ser (tcc); Val (gtc); and Arg (agg). All codons which do not fit the description of preferred codons or less preferred codons are non-preferred codons. In general, the degree of preference of a particular codon is indicated by the prevalence of the codon in highly expressed human genes as indicated in Table 1 under the heading "High." For example,

"atc" represents 77% of the Ile codons in highly expressed mammalian genes and is the preferred Ile codon; "att" represents 18% of the Ile codons in highly expressed mammalian genes and is the less preferred Ile codon. The sequence "ata" represents only 5% of the Ile codons in highly expressed human genes as is a non-preferred Ile codon. Replacing a codon with another codon that is more prevalent in highly expressed human genes will generally increase expression of the gene in mammalian cells. Accordingly, the invention includes replacing a less preferred codon with a preferred codon as well as replacing a non-preferred codon with a preferred or less preferred codon.

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By "protein normally expressed in a mammalian cell" is meant a protein which is expressed in mammalian under natural conditions. The term includes genes in the mammalian genome such as those encoding Factor VIII, Factor IX, interleukins, and other proteins. The term also includes genes which are expressed in a mammalian cell under disease conditions such as oncogenes as well as genes which are encoded by a virus (including a retrovirus) which are expressed in mammalian cells post-infection. By "protein normally expressed in a eukaryotic cell" is meant a protein which is expressed in a eukaryote under natural conditions. The term also includes genes which are expressed in a mammalian cell under disease conditions.

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In preferred embodiments, the synthetic gene is capable of expressing the mammalian or eukaryotic protein at a level which is at least 110%, 150%, 200%, 500%, 1,000%, 5,000% or even 10,000% of that expressed by the "natural" (or "native") gene in an *in vitro* mammalian cell culture system under identical conditions (i.e., same cell type, same culture conditions, same expression vector).

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Suitable cell culture systems for measuring expression of the synthetic gene and corresponding natural gene are described below. Other

suitable expression systems employing mammalian cells are well known to those skilled in the art and are described in, for example, the standard molecular biology reference works noted below. Vectors suitable for expressing the synthetic and natural genes are described below and in the standard reference works described below. By "expression" is meant protein expression. Expression can be measured using an antibody specific for the protein of interest. Such antibodies and measurement techniques are well known to those skilled in the art. By "natural gene" and "native gene" is meant the gene sequence (including naturally occurring allelic variants) which naturally encodes the protein, i.e., the native or natural coding sequence.

In other preferred embodiments at least 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, or 90% of the codons in the natural gene are non-preferred codons.

In other preferred embodiments at least 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, or 90% of the non-preferred codons in the natural gene are replaced with preferred codons or less preferred codons.

In other preferred embodiments at least 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, or 90% of the non-preferred codons in the natural genc are replaced with preferred codons.

In a preferred embodiment the protein is a retroviral protein. In a more preferred embodiment the protein is a lentiviral protein. In an even more preferred embodiment the protein is an HIV protein. In other preferred embodiments the protein is gag, pol, env, gp120, or gp160. In other preferred embodiments the protein is a human protein. In more preferred embodiments, the protein is human Factor VIII and the protein in B region deleted human Factor VIII. In another preferred embodiment the protein is green flourescent protein.

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In various preferred embodiments at least 30%, 40%, 50%, 60%, 70%, 80%, 90%, and 95% of the codons in the synthetic gene are preferred or less preferred codons.

The invention also features an expression vector comprising the synthetic gene.

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In another aspect the invention features a cell harboring the synthetic gene. In various preferred embodiments the cell is a prokaryotic cell and the cell is a mammalian cell.

In preferred embodiments the synthetic gene includes fewer than 50, fewer than 40, fewer than 30, fewer than 20, fewer than 10, fewer than 5, or no "cg" sequences.

The invention also features a method for preparing a synthetic gene encoding a protein normally expressed by a mammalian cell or other eukaryotic cell. The method includes identifying non-preferred and less-preferred codons in the natural gene encoding the protein and replacing one or more of the non-preferred and less-preferred codons with a preferred codon encoding the same amino acid as the replaced codon.

Under some circumstances (e.g., to permit introduction of a restriction site) it may be desirable to replace a non-preferred codon with a less preferred codon rather than a preferred codon.

It is not necessary to replace all less preferred or non-preferred codons with preferred codons. Increased expression can be accomplished even with partial replacement of less preferred or non-preferred codons with preferred codons. Under some circumstances it may be desirable to only partially replace non-preferred codons with preferred or less preferred codons in order to obtain an intermediate level of expression.

In other preferred embodiments the invention features vectors (including expression vectors) comprising one or more the synthetic genes.

By "vector" is meant a DNA molecule, derived, e.g., from a plasmid, bacteriophage, or mammalian or insect virus, into which fragments of DNA may be inserted or cloned. A vector will contain one or more unique restriction sites and may be capable of autonomous replication in a defined host or vehicle organism such that the cloned sequence is reproducible. Thus, by "expression vector" is meant any autonomous element capable of directing the synthesis of a protein. Such DNA expression vectors include mammalian plasmids and viruses.

The invention also features synthetic gene fragments which encode a desired portion of the protein. Such synthetic gene fragments are similar to the synthetic genes of the invention except that they encode only a portion of the protein. Such gene fragments preferably encode at least 50, 100, 150, or 500 contiguous amino acids of the protein.

In constructing the synthetic genes of the invention it may be desirable to avoid CpG sequences as these sequences may cause gene silencing. Thus, in a preferred embodiment the coding region of the synthetic gene does not include the sequence "cg."

The codon bias present in the HIV gp120 env gene is also present in the gag and pol genes. Thus, replacement of a portion of the non-preferred and less preferred codons found in these genes with preferred codons should produce a gene capable of higher level expression. A large fraction of the codons in the human genes encoding Factor VIII and Factor IX are non-preferred codons or less preferred codons. Replacement of a portion of these codons with preferred codons should yield genes capable of higher level expression in mammalian cell culture.

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The synthetic genes of the invention can be introduced into the cells of a living organism. For example, vectors (viral or non-viral) can be used to introduce a synthetic gene into cells of a living organism for gene therapy.

Conversely, it may be desirable to replace preferred codons in a naturally occurring gene with less-preferred codons as a means of lowering expression.

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Standard reference works describing the general principles of recombinant DNA technology include Watson et al., Molecular Biology of the Gene, Volumes I and II, the Benjamin/Cummings Publishing Company, Inc., publisher, Menlo Park, CA (1987); Darnell et al., Molecular Cell Biology, Scientific American Books, Inc., Publisher, New York, N.Y. (1986); Old et al., Principles of Gene Manipulation: An Introduction to Genetic Engineering, 2d edition, University of California Press, publisher, Berkeley, CA (1981); Maniatis et al., Molecular Cloning: A Laboratory Manual, 2nd Ed. Cold Spring Harbor Laboratory, publisher, Cold Spring Harbor, NY (1989); and Current Protocols in Molecular Biology, Ausubel et al., Wiley Press, New York, NY (1992).

By "transformed cell" is meant a cell into which (or into an ancestor of which) has been introduced, by means of recombinant DNA techniques, a selected DNA molecule, e.g., a synthetic gene.

By "positioned for expression" is meant that a DNA molecule, e.g., a synthetic gene, is positioned adjacent to a DNA sequence which directs transcription and translation of the sequence (i.e., facilitates the production of the protein encoded by the synthetic gene.

Description of the Drawings

Figure 1 depicts the sequence of the synthetic gp120 and a synthetic gp160 gene in which codons have been replaced by those found in highly expressed human genes.

Figure 2 is a schematic drawing of the synthetic gp120 (HIV-1 MN) gene. The shaded portions marked v1 to v5 indicate hypervariable regions. The filled box indicates the CD4 binding site. A limited number of the unique restriction sites ares shown: H (Hind3), Nh (Nhe1), P (Pst1), Na (Nae1), M (Mlu1), R (EcoR1), A (Age1) and No (Not1). The chemically synthesized DNA fragments which served as PCR templates are shown below the gp120 sequence, along with the locations of the primers used for their amplification.

Figure 3 is a photograph of the results of transient transfection assays used to measure gp120 expression. Gel electrophoresis of immunoprecipitated supernatants of 293T cells transfected with plasmids expressing gp120 encoded by the IIIB isolate of HIV-1 (gp120IIIb), by the MN isolate of HIV-1 (gp120mn), by the MN isolate of HIV-1 modified by substitution of the endogenous leader peptide with that of the CD5 antigen (gp120mnCD5L), or by the chemically synthesized gene encoding the MN variant of HIV-1 with the human CD5Leader (syngp120mn). Supernatants were harvested following a 12 hour labeling period 60 hours post-transfection and immunoprecipitated with CD4:IgG1 fusion protein and protein A sepharose.

Figure 4 is a graph depicting the results of ELISA assays used to measure protein levels in supernatants of transiently transfected 293T cells. Supernatants of 293T cells transfected with plasmids expressing gp120 encoded by the IIIB isolate of HIV-1 (gp120 IIIb), by the MN isolate of HIV-1 (gp120mn), by the MN isolate of HIV-1 modified by substitution of the endogenous leader peptide with that of CD5 antigen (gp120mn CD5L), or by

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the chemically synthesized gene encoding the MN variant of HIV-1 with human CDS leader (syngp120mn) were harvested after 4 days and tested in a gp120/CD4 ELISA. The level of gp120 is expressed in ng/ml.

Figure 5A is a photograph of a gel illustrating the results of a immunoprecipitation assay used to measure expression of the native and synthetic gp120 in the presence of rev in trans and the RRE in cis. In this experiment 293T cells were transiently transfected by calcium phosphate coprecipitation of 10 μ g of plasmid expressing: (A) the synthetic gp120MN sequence and RRE in cis, (B) the gp120 portion of HIV-1 IIIB, (C) the gp120 portion of HIV-1 IIIB and RRE in cis, all in the presence or absence of rev expression. The RRE constructs gp120IIIbRRE and syngp120mnRRE were generated using an Eag1/Hpa1 RRE fragment cloned by PCR from a HIV-1 HXB2 proviral clone. Each gp120 expression plasmid was cotransfected with 10 μ g of either pCMVrev or CDM7 plasmid DNA. Supernatants were harvested 60 hours post transfection, immunoprecipitated with CD4:IgG fusion protein and protein A agarose, and run on a 7% reducing SDS-PAGE. The gel exposure time was extended to allow the induction of gp120IIIbrre by rev to be demonstrated.

z Figure 5B is a shorter exposure of a similar experiment in which syngp120mnrre was cotransfected with or without pCMVrev.

Figure 5C is a schematic diagram of the constructs used in Figure 5A.

Figure 6 is a comparison of the sequence of the wild-type ratTHY-1 gene (wt) and a synthetic ratTHY-1 gene (env) constructed by chemical synthesis and having the most prevalent codons found in the HIV-1 env gene.

Figure 7 is a schematic diagram of the synthetic ratTHY-1 gene. The solid black box denotes the signal peptide. The shaded box denotes the

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sequences in the precursor which direct the attachment of a phophatidyl-inositol glycan anchor. Unique restriction sites used for assembly of the THY-1 constructs are marked H (Hind3), M (Mlu1), S (Sac1) and No (Not1). The position of the synthetic oligonucleotides employed in the construction are shown at the bottom of the figure.

Figure 8 is a graph depicting the results of flow cytometry analysis. In this experiment 293T cells transiently transfected with either a wild-type ratTHY-1 expression plasmid (thick line), ratTHY-1 with envelope codons expression plasmid (thin line), or vector only (dotted line) by calcium phosphate co-precipitation. Cells were stained with anti-ratTHY-1 monoclonal antibody OX7 followed by a polyclonal FITC-conjugated anti-mouse IgG antibody 3 days after transfection.

Figure 9A is a photograph of a gel illustrating the results of immunoprecipitation analysis of supernatants of human 293T cells transfected with either syngp120mn (A) or a construct syngp120mn.rTHY-1env which has the rTHY-1env gene in the 3' untranslated region of the syngp120mn gene (B). The syngp120mn.rTHY-1env construct was generated by inserting a Not1 adapter into the blunted Hind3 site of the rTHY-1env plasmid. Subsequently, a 0.5 kb Not1 fragment containing the rTHY-1env gene was cloned into the Not1 site of the syngp120mn plasmid and tested for correct orientation. Supernatants of 35S labeled cells were harvested 72 hours post transfection, precipitated with CD4:IgG fusion protein and protein A agarose, and run on a 7% reducing SDS-PAGE.

Figure 9B is a schematic diagram of the constructs used in the experiment depicted in Figure 9A.

Figure 10A is a photograph of COS cells transfected with vector only showing no GFP fluorescence.

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Figure 10B is a photograph of COS cells transfected with a CDM7 expression plasmid encoding native GFP engineered to include a consensus translational initiation sequence.

Figure 10C is a photograph of COS cells transfected with an expression plasmid having the same flanking sequences and initiation consensus as in Figure 10B, but bearing a codon optimized gene sequence.

Figure 10D is a photograph of COS cells transfected with an expression plasmid as in Figure 10C, but bearing a Thr at residue 65 in place of Ser.

Figure 11 depicts the sequence of a synthetic gene encoding green flourescent proteins (SEQ ID NO:40).

Figure 12 depicts the sequence of a native human Factor VIII gene lacking the central B domain (amino acids 760-1639, inclusive) (SEQ ID NO:41).

Figure 13 depicts the sequence of a synthetic human Factor VIII gene lacking the central B domain (amino acids 760-1639, inclusive) (SEQ ID NO:42).

Description of the Preferred Embodiments

EXAMPLE 1

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20 Construction of a Synthetic gp120 Gene Having Codons Found in Highly
Expressed-Human-Genes

A codon frequency table for the envelope precursor of the LAV subtype of HIV-1 was generated using software developed by the University of Wisconsin Genetics Computer Group. The results of that tabulation are contrasted in Table 1 with the pattern of codon usage by a collection of highly expressed human genes. For any amino acid encoded by degenerate codons,

the most favored codon of the highly expressed genes is different from the most favored codon of the HIV envelope precursor. Moreover a simple rule describes the pattern of favored envelope codons wherever it applies: preferred codons maximize the number of

adenine residues in the viral RNA. In all cases but one this means that the codon in which the third position is A is the most frequently used. In the special case of serine, three codons equally contribute one A residue to the mRNA; together these three comprise 85% of the serine codons actually used in envelope transcripts. A particularly striking example of the A bias is found in the codon choice for arginine, in which the AGA triplet comprises 88% of the arginine codons. In addition to the preponderance of A residues, a marked preference is seen for uridine among degenerate codons whose third residue must be a pyrimidine. Finally, the inconsistencies among the less frequently used variants can be accounted for by the observation that the dinucleotide CpG is under represented; thus the third position is less likely to be G whenever the second position is C, as in the codons for alanine, proline, serine and threonine; and the CGX triplets for arginine are hardly used at all.

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TABLE 1:	Codon Frequency in the I	HIV-1 IIIb env	v gene and in highl	v
	expressed human genes.	,	, , ,	_

			High	Env	• .		٠.,		. 4	High	Env
	Ala			ı	,			Cys		, 0	
5	GC	\mathbf{C}	53	27		,	,	TG	С	68	16 ,
		T	, 17	18		1	' · ·	, , , ,	T ·	32	84
		Α		50			٠.,	The second	_		* .
		G '	17	5				<u>Gln</u>		1	1
	• •			1 7				CA.	A	12	55
10	Arg							1	G	88	45
	CG	C	37	0			. ,	i i	J	00	1
		T '	7	Y.A				Glu			
		Ā	6	0	•		i	GA	Α	25	67
		G G	21	0			•	UA	G	75	
15	AG	A	10	88					· U	13	33
13	AU	G'	18	8				Ch.	٠.,		t .
		u	10	0			1	Gly		50	
	Aom	٠.,				,		GG	C	50	6
	Asn A A	C	70	120			•		T	12	13
20	AA	C	78 22	30					A	14	53
20		T	22	70					G	24	28
	Asp							<u>His</u>			
	GA	C	75	33			•	CA	С	79	25
		T	25	67					T	21	75
	. *	-							•		.5
								<u>Ile</u>			
25								AT	C	77	25
	•	,							T	18	31
									Α	5	44
	<u>Leu</u>		1		_			Sor			
	CT	С	26	10				Ser	C	20	O
30	CI	T		7.				TC	C	28	8
30			5 3						T	13	8
		A		17				•	A	5	22
	TT	G	58	17					G	9	0
	TT	A	2	30				AG	C	34	22
		G	6	20					T	10	41

	Lys					1,	Thr		,	l
	AA	ļΑ	18	68	1		AC .	C ,	57 ·	20
		, G	82	32 ;	1			T ' '	14	22
		1	•	t el		•		\mathbf{A}^{-1}	14	51
5					•		, ,	\mathbf{G}	15	7
		100	1			1 ,				4
	<u>Pro</u>	, ,		1	1		Tyr	1		
	CC	¹ C	48	27		٠	TA	Ç	74.	8
•		\mathbf{T}^{-1}	19	14		•		T	26	92
	•	Α	16.	· ·55			1		•	
10		G	17	5			1			
							1			1 '
	<u>Phe</u>	1	•	.Ж.			<u>Yal</u>			
	TT	\mathbf{C}^{-1}	80 ,	26	•		GT	C	25	12
		T	20	74		1.		T	7	9
			1	I				A	5	62
15						1	i	G	64	18

Codon frequency was calculated using the GCG program established the University of Wisconsin Genetics Computer Group. Numbers represent the percentage of cases in which the particular codon is used. Codon usage frequencies of envelope genes of other HIV-1 virus isolates are comparable and show a similar bias.

In order to produce a gp120 gene capable of high level expression in mammalian cells, a synthetic gene encoding the gp120 segment of HIV-1 was constructed (syngp120mn), based on the sequence of the most common North American subtype, HIV-1 MN (Shaw et al., Science 226:1165, 1984; Gallo et al., Nature 321:119, 1986). In this synthetic gp120 gene nearly all of the native codons have been systematically replaced with codons most frequently used in highly expressed human genes (Figure 1). This synthetic gene was assembled from chemically synthesized oligonucleotides of 150 to 200 bases in length. If oligonucleotides exceeding 120 to 150 bases are chemically synthesized, the

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percentage of full-length product can be low, and the vast excess of material consists of shorter oligonucleotides. Since these shorter fragments inhibit cloning and PCR procedures, it can be very difficult to use oligonucleotides exceeding a certain length. In order to use crude synthesis material without prior purification, single-stranded oligonucleotide pools were PCR amplified before cloning. PCR products were purified in agarose gels and used as templates in the next PCR step. Two adjacent fragments could be co-amplified because of overlapping sequences at the end of either fragment. These fragments, which were between 350 and 400 bp in size, were subcloned into a pCDM7-derived plasmid containing the leader sequence of the CD5 surface molecule followed by a Nhe1/Pst1/Mlu1/EcoR1/BamH1 polylinker. Each of the restriction enzymes in this polylinker represents a site that is present at either the 5' or 3' end of the PCR-generated fragments. Thus, by sequential subcloning of each of the 4 long fragments, the whole gp120 gene was assembled. For each fragment three to six different clones were subcloned and sequenced prior to assembly. A schematic drawing of the method used to construct the synthetic gp120 is shown in Figure 2. The sequence of the synthetic gp120 gene (and a synthetic gp160 gene created using the same approach) is presented in Figure 1.

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The mutation rate was considerable. The most commonly found mutations were short (1 nucleotide) and long (up to 30 nucleotides) deletions.

In some cases it was necessary to exchange parts with either synthetic adapters

or pieces from other subclones without mutation in that particular region.

Some deviations from strict adherence to optimized codon usage were made to accommodate the introduction of restriction sites into the resulting gene to facilitate the replacement of various segments (Figure 2). These unique restriction sites were introduced into the gene at approximately 100 bp

intervals. The native HIV leader sequence was exchanged with the highly efficient leader peptide of the human CD5 antigen to facilitate secretion (Aruffo et al., Cell 61:1303, 1990) The plasmid used for construction is a derivative of the mammalian expression vector pCDM7 transcribing the inserted gene under the control of a strong human CMV immediate early promoter.

To compare the wild-type and synthetic gp120 coding sequences, the synthetic gp120 coding sequence was inserted into a mammalian expression vector and tested in transient transfection assays. Several different native gp120 genes were used as controls to exclude variations in expression levels between different virus isolates and artifacts induced by distinct leader sequences. The gp120 HIV IIIb construct used as control was generated by PCR using a Sal1/Xho1 HIV-1 HXB2 envelope fragment as template. To exclude PCR induced mutations, a Kpn1/Ear1 fragment containing approximately 1.2 kb of the gene was exchanged with the respective sequence from the proviral clone. The wild-type gp120mn constructs used as controls were cloned by PCR from HIV-1 MN infected C8166 cells (AIDS Repository, Rockville, MD) and expressed gp120 either with a native envelope or a CD5 leader sequence. Since proviral clones were not available in this case, two clones of each construct were tested to avoid PCR artifacts. To determine the amount of secreted gp120 semi-quantitatively supernatants of 293T cells transiently transfected by calcium phosphate co-precipitation were immunoprecipitated with soluble CD4:immunoglobulin fusion protein and protein A sepharose.

The results of this analysis (Figure 3) show that the synthetic gene product is expressed at a very high level compared to that of the native gp120 controls. The molecular weight of the synthetic gp120 gene was comparable to

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control proteins (Figure 3) and appeared to be in the range of 100 to 110 kd. The slightly faster migration can be explained by the fact that in some tumor cell lines, e.g., 293T, glycosylation is either not complete or altered to some extent.

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To compare expression more accurately gp120 protein levels were quantitated using a gp120 ELISA with CD4 in the demobilized phase. This analysis shows (Figure 4) that ELISA data were comparable to the immunoprecipitation data, with a gp120 concentration of approximately 125 ng/ml for the synthetic gp120 gene, and less than the background cutoff (5 ng/ml) for all the native gp120 genes. Thus, expression of the synthetic gp120 gene appears to be at least one order of magnitude higher than wild-type gp120 genes. In the experiment shown the increase was at least 25 fold.

The Role of rev in gp120 Expression

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of a viral transcript, the possible role of non-translational effects in the improved expression of the synthetic gp120 gene was tested. First, to rule out the possibility that negative signals elements conferring either increased mRNA degradation or nucleic retention were eliminated by changing the nucleotide sequence, cytoplasmic mRNA levels were tested. Cytoplasmic RNA was prepared by NP40 lysis of transiently transfected 293T cells and subsequent elimination of the nuclei by centrifugation. Cytoplasmic RNA was

Since rev appears to exert its effect at several steps in the expression

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subsequently prepared from lysates by multiple phenol extractions and precipitation, spotted on nitrocellulose using a slot blot apparatus, and finally hybridized with an envelope-specific probe.

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Briefly, cytoplasmic mRNA 293 cells transfected with CDM&, gp120 IIIB, or syngp120 was isolated 36 hours post transfection. Cytoplasmic RNA of Hela cells infected with wild-type vaccinia virus or recombinant virus

expressing gp120 IIIb or the synthetic gp120 gene was under the control of the 7.5 promoter was isolated 16 hours post infection. Equal amounts were spotted on nitrocellulose using a slot blot device and hybridized with randomly labeled 1.5 kb gp120IIIb and syngp120 fragments or human beta-actin. RNA expression levels were quantitated by scanning the hybridized membranes with a phospoimager. The procedures used are described in greater detail below.

This experiment demonstrated that there was no significant difference in the mRNA levels of cells transfected with either the native or synthetic gp120 gene. In fact, in some experiments cytoplasmic mRNA level of the synthetic gp120 gene was even lower than that of the native gp120 gene.

These data were confirmed by measuring expression from recombinant vaccinia viruses. Human 293 cells or Hela cells were infected with vaccinia virus expressing wild-type gp120 IIIb or syngp120mn at a multiplicity of infection of at least 10. Supernatants were harvested 24 hours post infection and immunoprecipitated with CD4:immunoglobin fusion protein and protein A sepharose. The procedures used in this experiment are described in greater detail below.

This experiment showed that the increased expression of the synthetic gene was still observed when the endogenous gene product and the synthetic gene product were expressed from vaccinia virus recombinants under the control of the strong mixed early and late 7.5k promoter. Because vaccinia virus mRNAs are transcribed and translated in the cytoplasm, increased expression of the synthetic envelope gene in this experiment cannot be attributed to improved export from the nucleus. This experiment was repeated in two additional human cell types, the kidney cancer cell line 293 and HeLa cells. As with transfected 293T cells, mRNA levels were similar in 293 cells infected with either recombinant vaccinia virus.

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Codon Usage in Lentivirus

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Because it appears that codon usage has a significant impact on expression in mammalian cells, the codon frequency in the envelope genes of other retroviruses was examined. This study found no clear pattern of codon preference between retroviruses in general. However, if viruses from the lentivirus genus, to which HIV-1 belongs to, were analyzed separately, codon usage bias almost identical to that of HIV-1 was found. A codon frequency table from the envelope glycoproteins of a variety of (predominantly type C) retroviruses excluding the lentiviruses was prepared, and compared a codon frequency table created from the envelope sequences of four lentiviruses not closely related to HIV-1 (caprine arthritis encephalitis virus, equine infectious ancmia virus, feline immunodeficiency virus, and visna virus) (Table 2). The codon usage pattern for lentiviruses is strikingly similar to that of HIV-1, in all cases but one, the preferred codon for HIV-1 is the same as the preferred codon for the other lentiviruses. The exception is proline, which is encoded by CCT in 41% of non-HIV lentiviral envelope residues, and by CCA in 40% of residues, a situation which clearly also reflects a significant preference for the triplet ending in A. The pattern of codon usage by the non-lentiviral envelope proteins does not show a similar predominance of A residues, and is also not as skewed toward third position C and G residues as is the codon usage for the highly expressed human genes. In general non-lentiviral retroviruses appear to exploit the different codons more equally, a pattern they share with less highly expressed human genes.

TABLE 2: Codon frequency in the envelope gene of lentiviruses (lenti) and non-lentiviral retroviruses (other)

		' C	ther L	enti .			O	Other Lenti				
	<u>Ala</u>			1	1		Cys	,	1			
5	GC	· C , i	45	13			TG	C	53	21,		
	•	' T '	26	.37		· · · .	, , ,	T.	47	79		
		1 A	20	46	,	1	· ('			٠.		
		\mathbf{G}^{-1}		3			<u>Gln</u>	1 1	1	1		
	•						CA	Α	52	69		
10	Arg	•					1	G	48	31		
	CG	С	.14	' 2			1	J	10			
		T '	6	× 3			<u>Glu</u>					
		Ā	16	11 5 5	•		GA	· A	57	68 -		
		G	17	3			, U A	G	43	32		
15	AG	A	3,1	' 5 1	•				43	32		
	110	G	15	26			Gly	٠.		,		
		J	1,5	20		,	GG	С	21	ο.		
	Asn	• .					00	T		8		
	AA	С	49	31		,			13	• 9		
20	лл	T						A	37.	56		
20		1	51	69				G	29	26		
	Asp						His					
	GA	С	55	33		•	CA	С	51	38		
		T	51	69				Т	49	62		
							<u>Ile</u>					
25							AT	C	38	16		
								T	31	22		
								Α	31	61		
	Lau		•				C		i			
	<u>Leu</u> CT	С	22	8		-	<u>Ser</u>	~	20	10		
2.0	CI		22				TC	C	38	10		
30		T	14	9				T	17	16		
		A	21	16				A	18	24		
	TT	G	19	11			. ~	G	6	5		
	TT	A	15	41			AG	C	13	20		
		G	10	16				T	7	25		

					4						'
						·				•	
	Lys				•	•	Thr.	ŀ			
•	ĄΑ	Α	60	63	* 4 ₁		AC	, C	44	18	
	. ,	' G	40	37	ı		-X	T	' 27	1 20	
					•	. 1		Α	19	55	
5	Pro							G	10	8	
	CC	С	42	14	1						
	•	· T	30	41		+	Tyr				,
	1.	' A	20	40	ı		TA	C	48	28	
		· G ·	7	5,				T .	52	72	
10	<u>Phe</u>						<u>Val</u>				
ı	TŢ	\mathbf{C}	52	25			GT	C	36	9	
	,	T	48	75				T.	-17	10	
							•	Α	22	54	
				: '				G	25	27	
15		•								l	

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Codon frequency was calculated using the GCG program established by the University of Wisconsin Genetics Computer Group. Numbers represent the percentage in which a particular codon is used. Codon usage of non-lentiviral retroviruses was compiled from the envelope precursor sequences of bovine leukemia virus feline leukemia virus, human T-cell leukemia virus type I, human T-cell lymphotropic virus type II, the mink cell focus-forming isolate of murine leukemia virus (MuLV), the Rauscher spleen focus-forming isolate, the 10A1 isolate, the 4070A amphotropic isolate and the myeloproliferative leukemia virus isolate, and from rat leukemia virus, simian sarcoma virus, simian T-cell leukemia virus, leukemogenic retrovirus T1223/B and gibbon ape 25. leukemia virus. The codon frequency tables for the non-HIV, non-SIV lentiviruses were compiled from the envelope precursor sequences for caprine arthritis encephalitis virus, equine infectious anemia virus, feline immunodeficiency virus, and visna virus.

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In addition to the prevalence of codons containing an A, lentiviral codons adhere to the HIV pattern of strong CpG under representation, so that the third position for alanine, proline, serine and threonine triplets is rarely G. The retroviral envelope triplets show a similar, but less pronounced, under representation of CpG. The most obvious difference between lentiviruses and

other retroviruses with respect to CpG prevalence lies in the usage of the CGX variant of arginine triplets, which is reasonably frequently represented among the retroviral envelope coding sequences, but is almost never present among the comparable lentivirus sequences.

Differences in rev Dependence Between Native and Synthetic gp120

To examine whether regulation by rev is connected to HIV-1 codon usage, the influence of rev on the expression of both native and synthetic gene was investigated. Since regulation by rev requires the rev-binding site RRE in cis, constructs were made in which this binding site was cloned into the 3' untranslated region of both the native and the synthetic gene. These plasmids were co-transfected with rev or a control plasmid in trans into 293T cells, and gp120 expression levels in supernatants were measured semiquantitatively by immunoprecipitation. The procedures used in this experiment are described in greater detail below.

As shown in Figure 5A and Figure 5B, rev up regulates the native gp120 gene, but has no effect on the expression of the synthetic gp120 gene. Thus, the action of rev is not apparent on a substrate which lacks the coding sequence of endogenous viral envelope sequences.

Expression of a synthetic ratTHY-1 gene with HIV envelope codons

The above-described experiment suggest that in fact "envelope sequences" have to be present for rev regulation. In order to test this hypothesis, a synthetic version of the gene encoding the small, typically highly expressed cell surface protein, ratTHY-1 antigen, was prepared. The synthetic version of the ratTHY-1 gene was designed to have a codon usage like that of HIV gp120. In designing this synthetic gene AUUUA sequences, which are associated with mRNA instability, were avoided. In addition, two restriction

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sites were introduced to simplify manipulation of the resulting gene (Figure 6). This synthetic gene with the HIV envelope codon usage (rTHY-1env) was generated using three 150 to 170 mer oligonucleotides (Figure 7). In contrast to the syngp120mn gene, PCR products were directly cloned and assembled in pUC12, and subsequently cloned into pCDM7.

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Expression levels of native rTHY-1 and rTHY-1 with the HIV envelope codons were quantitated by immunofluorescence of transiently transfected 293T cells. Figure 8 shows that the expression of the native THY-1 gene is almost two orders of magnitude above the background level of the control transfected cells (pCDM7). In contrast, expression of the synthetic ratTHY-1 is substantially lower than that of the native gene (shown by the shift to of the peak towards a lower channel number).

To prove that no negative sequence elements promoting mRNA degradation were inadvertently introduced, a construct was generated in which the rTHY-1env gene was cloned at the 3' end of the synthetic gp120 gene (Figure 9B). In this experiment 293T cells were transfected with either the syngp120mn gene or the syngp120/ratTHY-1 env fusion gene (syngp120mn.rTHY-1env). Expression was measured by immunoprecipitation with CD4:IgG fusion protein and protein A agarose. The procedures used in this experiment are described in greater detail below.

Since the synthetic gp120 gene has an UAG stop codon, rTHY-1env is not translated from this transcript. If negative elements conferring enhanced degradation were present in the sequence, gp120 protein levels expressed from this construct should be decreased in comparison to the syngp120mn construct without rTHY-1env. Figure 9A, shows that the expression of both constructs is similar, indicating that the low expression must be linked to translation.

Rev-dependent expression of synthetic ratTHY-1 gene with envelope codons

To explore whether rev is able to regulate expression of a ratTHY-1 gene having env codons, a construct was made with a rev-binding site in the 3' end of the rTHY1env open reading frame. To measure rev-responsiveness of the a ratTHY-1env construct having a 3' RRE, human 293T cells were cotransfected ratTHY-1envrrc and either CDM7 or pCMVrev. At 60 hours post transfection cells were detached with 1 mM EDTA in PBS and stained with the OX-7 anti rTHY-1 mouse monoclonal antibody and a secondary FITC-conjugated antibody. Fluorescence intensity was measured using a EPICS XL cytofluorometer. These procedures are described in greater detail below.

In repeated experiments, a slight increase of rTHY-1env expression was detected if rev was cotransfected with the rTHY-1env gene. To further increase the sensitivity of the assay system a construct expressing a secreted version of rTHY-1env was generated. This construct should produce more reliable data because the accumulated amount of secreted protein in the supernatant reflects the result of protein production over an extended period, in contrast to surface expressed protein, which appears to more closely reflect the current production rate. A gene capable of expressing a secreted form was prepared by PCR using forward and reverse primers annealing 3' of the endogenous leader sequence and 5' of the sequence motif required for phosphatidylinositol glycan anchorage respectively. The PCR product was cloned into a plasmid which already contained a CD5 leader sequence, thus generating a construct in which the membrane anchor has been deleted and the leader sequence exchanged by a heterologous (and probably more efficient) leader peptide.

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The rev-responsiveness of the secreted form ratTHY-1env was measured by immunoprecipitation of supernatants of human 293T cells cotransfected with a plasmid expressing a secreted form of ratTHY-1env and the RRE sequence in cis (rTHY-1envPI-rre) and either CDM7 or pCMVrev. The rTHY-1envPI-RRE construct was made by PCR using the oligonucleotide: cgcggggctagcgcaaagagtaataagtttaac (SEQ ID NO:38) as a forward primer, the oligonucleotide: cgcggatcccttgtattttgtactaata (SEQ ID NO:39) as reverse primer, and the synthetic rTHY-1env construct as a template. After digestion with Nhe1 and Not1 the PCR fragment was cloned into a plasmid containing CD5 leader and RRE sequences. Supernatants of ³⁵S labeled cells were harvested 72 hours post transfection, precipitated with a mouse monoclonal antibody OX7 against rTHY-1 and anti mouse IgG sepharose, and run on a 12% reducing SDS-PAGE.

In this experiment the induction of rTHY-1env by rev was much more prominent and clear-cut than in the above-described experiment and strongly suggests that rev is able to translationally regulate transcripts that are suppressed by low-usage codons.

Rev-independent expression of a rTHY-lenv:immunoglobulin fusion protein

To test whether low-usage codons must be present throughout the whole coding sequence or whether a short region is sufficient to confer revresponsiveness, a rTHY-lenv:immunoglobulin fusion protein was generated. In this construct the rTHY-lenv genc (without the sequence motif responsible for phosphatidylinositol glycan anchorage) is linked to the human IgG1 hinge, CH2 and CH3 domains. This construct was generated by anchor PCR using primers with Nhe1 and BamHI restriction sites and rTHY-lenv as template. The PCR fragment was cloned into a plasmid containing the leader sequence of

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the CD5 surface molecule and the hinge, CH2 and CH3 parts of human IgG1 immunoglobulin. A Hind3/Eag1 fragment containing the rTHY-1enveg1 insert was subsequently cloned into a pCDM7-derived plasmid with the RRE sequence.

To measure the response of the rTHY-1env/ immunoglobin fusion gene (rTHY-1enveg1rre) to rev human 293T cells cotransfected with rTHY-1enveg1rre and either pCDM7 or pCMVrev. The rTHY-1enveg1rre construct was made by anchor PCR using forward and reverse primers with Nhe1 and BamH1 restriction sites respectively. The PCR fragment was cloned into a plasmid containing a CD5 leader and human IgG1 hinge, CH2 and CH3 domains. Supernatants of ³⁵S labeled cells were harvested 72 hours post transfection, precipitated with a mouse monoclonal antibody OX7 against rTHY-1 and anti mouse IgG sepharose, and run on a 12% reducing SDS-PAGE. The procedures used are described in greater detail below.

As with the product of the rTHY-lenvPl- gene, this rTHY-lenv/immunoglobulin fusion protein is secreted into the supernatant. Thus, this gene should be responsive to rev-induction. However, in contrast to rTHY-lenvPl-, cotransfection of rev in trans induced no or only a negligible increase of rTHY-lenvegl expression.

The expression of rTHY-1:immunoglobulin fusion protein with native rTHY-1 or HIV envelope codons was measured by immunoprecipitation. Briefly, human 293T cells transfected with either rTHY-1enveg1 (env codons) or rTHY-1wteg1 (native codons). The rTHY-1wteg1 construct was generated in manner similar to that used for the rTHY-1enveg1 construct, with the exception that a plasmid containing the native rTHY-1 gene was used as template. Supernatants of ³⁵S labeled cells were harvested 72 hours post transfection, precipitated with a mouse monoclonal antibody OX7 against

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rTHY-1 and anti mouse IgG sepharose, and run on a 12% reducing SDS-PAGE. THE procedures used in this experiment are described in greater detail below.

Expression levels of rTHY-lenveg1 were decreased in comparison to a similar construct with wild-type rTHY-1 as the fusion partner, but were still considerably higher than rTHY-lenv. Accordingly, both parts of the fusion protein influenced expression levels. The addition of rTHY-lenv did not restrict expression to an equal level as seen for rTHY-lenv alone. Thus, regulation by rev appears to be ineffective if protein expression is not almost completely suppressed.

Codon preference in HIV-1 envelope genes

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Direct comparison between codon usage frequency of HIV envelope and highly expressed human genes reveals a striking difference for all twenty amino acids. One simple measure of the statistical significance of this codon preference is the finding that among the nine amino acids with two fold codon degeneracy, the favored third residue is A or U in all nine. The probability that all nine of two equiprobable choices will be the same is approximately 0.004, and hence by any conventional measure the third residue choice cannot be considered random. Further evidence of a skewed codon preference is found among the more degenerate codons, where a strong selection for triplets bearing adenine can be seen. This contrasts with the pattern for highly expressed genes, which favor codons bearing C, or less commonly G, in the third position of codons with three or more fold degeneracy.

The systematic exchange of native codons with codons of highly expressed human genes dramatically increased expression of gp120. A quantitative analysis by ELISA showed that expression of the synthetic gene was at least 25 fold higher in comparison to native gp120 after transient

transfection into human 293 cells. The concentration levels in the ELISA experiment shown were rather low. Since an ELISA was used for quantification which is based on gp120 binding to CD4, only native, non-denatured material was detected. This may explain the apparent low expression. Measurement of cytoplasmic mRNA levels demonstrated that the difference in protein expression is due to translational differences and not mRNA stability.

Retroviruses in general do not show a similar preference towards A and T as found for HIV. But if this family was divided into two subgroups, lentiviruses and non-lentiviral retroviruses, a similar preference to A and, less frequently, T, was detected at the third codon position for lentiviruses. Thus, the availing evidence suggests that lentiviruses retain a characteristic pattern of envelope codons not because of an inherent advantage to the reverse transcription or replication of such residues, but rather for some reason peculiar to the physiology of that class of viruses. The major difference between lentiviruses and non-complex retroviruses are additional regulatory and nonessentially accessory genes in lentiviruses, as already mentioned. Thus, one simple explanation for the restriction of envelope expression might be that an important regulatory mechanism of one of these additional molecules is based on it. In fact, it is known that one of these proteins, rev, which most likely has homologues in all lentiviruses. Thus codon usage in viral mRNA is used to create a class of transcripts which is susceptible to the stimulatory action of rev. This hypothesis was proved using a similar strategy as above, but this time codon usage was changed into the inverse direction. Codon usage of a highly expressed cellular gene was substituted with the most frequently used codons in the HIV envelope. As assumed, expression levels were considerably lower in comparison to the native molecule, almost two orders of magnitude when

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analyzed by immunofluorescence of the surface expressed molecule. If rev was coexpressed in trans and a RRE element was present in cis only a slight induction was found for the surface molecule. However, if THY-1 was expressed as a secreted molecule, the induction by rev was much more prominent, supporting the above hypothesis. This can probably be explained by accumulation of secreted protein in the supernatant, which considerably amplifies the rev effect. If rev only induces a minor increase for surface molecules in general, induction of HIV envelope by rev cannot have the purpose of an increased surface abundance, but rather of an increased intracellular gp160 level. It is completely unclear at the moment why this should be the case.

To test whether small subtotal elements of a gene are sufficient to restrict expression and render it rev-dependent rTHY1env:immunoglobulin fusion proteins were generated, in which only about one third of the total gene had the envelope codon usage. Expression levels of this construct were on an intermediate level, indicating that the rTHY-1env negative sequence element is not dominant over the immunoglobulin part. This fusion protein was not or only slightly rev-responsive, indicating that only genes almost completely suppressed can be rev-responsive.

Another characteristic feature that was found in the codon frequency tables is a striking under representation of CpG triplets. In a comparative study of codon usage in E. coli, yeast, drosophila and primates it was shown that in a high number of analyzed primate genes the 8 least used codons contain all codons with the CpG dinucleotide sequence. Avoidance of codons containing this dinucleotide motif was also found in the sequence of other retroviruses. It seems plausible that the reason for under representation of CpG-bearing triplets has something to do with avoidance of gene silencing by methylation of CpG

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cytosines. The expected number of CpG dinucleotides for HIV as a whole is about one fifth that expected on the basis of the base composition. This might indicate that the possibility of high expression is restored, and that the gene in fact has to be highly expressed at some point during viral pathogenesis.

The results presented herein clearly indicate that codon preference has a severe effect on protein levels, and suggest that translational elongation is controlling mammalian gene expression. However, other factors may play a role. First, abundance of not maximally loaded mRNA's in eukaryotic cells indicates that initiation is rate limiting for translation in at least some cases, since otherwise all transcripts would be completely covered by ribosomes. Furthermore, if ribosome stalling and subsequent mRNA degradation were the mechanism, suppression by rare codons could most likely not be reversed by any regulatory mechanism like the one presented herein. One possible explanation for the influence of both initiation and elongation on translational activity is that the rate of initiation, or access to ribosomes, is controlled in part by cues distributed throughout the RNA, such that the lentiviral codons predispose the RNA to accumulate in a pool of poorly initiated RNAs. However, this limitation need not be kinetic; for example, the choice of codons could influence the probability that a given translation product, once initiated, is properly completed. Under this mechanism, abundance of less favored codons would incur a significant cumulative probability of failure to complete the nascent polypeptide chain. The sequestered RNA would then be lent an improved rate of initiation by the action of rev. Since adenine residues are abundant in rev-responsive transcripts, it could be that RNA adenine methylation mediates this translational suppression.

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Detailed Procedures

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The following procedures were used in the above-described experiments.

Sequence Analysis

Sequence analyses employed the software developed by the University of Wisconsin Computer Group.

Plasmid constructions

Plasmid constructions employed the following methods. Vectors and insert DNA was digested at a concentration of 0.5 μ g/10 μ l in the appropriate restriction buffer for 1 - 4 hours (total reaction volume approximately 30 μ l). Digested vector was treated with 10% (v/v) of 1 μ g/ml calf intestine alkaline phosphatase for 30 min prior to gel electrophoresis. Both vector and insert digests (5 to 10 μ l each) were run on a 1.5% low melting agarose gel with TAE buffer. Gel slices containing bands of interest were transferred into a 1.5 ml reaction tube, melted at 65°C and directly added to the ligation without removal of the agarose. Ligations were typically done in a total volume of 25 μ l in 1x Low Buffer 1x Ligation Additions with 200-400 U of ligase, 1 μ l of vector, and 4 μ l of insert. When necessary, 5' overhanging ends were filled by ' adding 1/10 volume of 250 μ M dNTPs and 2-5 U of Klenow polymerase to heat inactivated or phenol extracted digests and incubating for approximately 20 min at room temperature. When necessary, 3' overhanging ends were filled by adding 1/10 volume of 2.5 mM dNTPs and 5-10 U of T4 DNA polymerase to heat inactivated or phenol extracted digests, followed by incubation at 37°C for 30 min. The following buffers were used in these reactions: 10x Low buffer (60 mM Tris HCl, pH 7.5, 60 mM MgCl₂, 50 mM NaCl, 4 mg/ml BSA, 70 mM β-mercaptoethanol, 0.02% NaN₃); 10x Medium buffer (60 mM Tris HCl, pH 7.5, 60 mM MgCl₂, 50 mM NaCl, 4 mg/ml BSA, 70 mM β-

mercaptoethanol, 0.02% NaN₃); 10x High buffer (60 mM Tris HCl, pH 7.5, 60 mM MgCl₂, 50 mM NaCl, 4 mg/ml BSA, 70 mM β-mercaptoethanol, 0.02% NaN₃); 10x Ligation additions (1 mM ATP, 20 mM DTT, 1 mg/ml BSA, 10 mM spermidine); 50x TAE (2 M Tris acetate, 50 mM EDTA).

Oligonucleotide synthesis and purification

Oligonucleotides were produced on a Milligen 8750 synthesizer (Millipore). The columns were eluted with 1 ml of 30% ammonium hydroxide, and the eluted oligonucleotides were deblocked at 55°C for 6 to 12 hours. After deblockiong, 150 μ l of oligonucleotide were precipitated with 10x volume of unsaturated n-butanol in 1.5 ml reaction tubes, followed by centrifugation at 15,000 rpm in a microfuge. The pellet was washed with 70% ethanol and resuspended in 50 μ l of H₂0. The concentration was determined by measuring the optical density at 260 nm in a dilution of 1:333 (1 OD₂₆₀ = 30 μ g/ml).

The following oligonucleotides were used for construction of the synthetic gp120 gene (all sequences shown in this text are in 5' to 3' direction).

oligo 1 forward (Nhe1): cgc ggg cta gcc acc gag aag ctg (SEQ ID NO:1).

oligo 1: acc gag aag ctg tgg gtg acc gtg tac tac ggc gtg ccc gtg tgg

aag ag ag gcc acc acc ctg ttc tgc gcc agc gac gcc aag gcg tac gac acc gag

gtg cac aac gtg tgg gcc acc cag gcg tgc gtg ccc acc gac ccc aac ccc cag gag gtg

gag ctc gtg aac gtg acc gag aac ttc aac at (SEQ ID NO:2).

oligo 1 reverse: cca cca tgt tgt tct tcc aca tgt tga agt tct c (SEQ ID NO:3).

oligo 2 forward: gac cga gaa ctt caa cat gtg gaa gaa caa cat (SEQ ID NO:4)

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oligo 2 reverse (Pst1): gtt gaa get gea gtt ett eat ete gee gee ett (SEQ ID NO:6).

oligo 3 forward (Pst1): gaa gaa ctg cag ctt caa cat cac cac cag c (SEQ ID NO:7).

oligo 3: aac atc acc acc agc atc cgc gac aag atg cag aag gag tac gcc ctg ctg tac aag ctg gat atc gtg agc atc gac aac gac agc acc agc tac cgc ctg atc tcc tgc aac acc agc gtg atc acc cag gcc tgc ccc aag atc agc ttc gag ccc atc ccc atc cac tac tgc gcc ccc gcc ggc ttc gcc (SEQ ID NO:8).

oligo 3 reverse: gaa ctt ctt gtc ggc ggc gaa gcc ggc ggg (SEQ ID NO:9).

oligo 4 forward: gcg ccc ccg ccg gct tcg cca tcc tga agt gca acg aca aga agt tc (SEQ ID NO:10)

oligo 4: gec gac aag aag tte age gge aag gge age tge aag aac gtg age ace gtg eag tge ace eac gge ate egg eeg gtg gtg age ace eag ete etg etg aac gge age etg gec gag gag gag gtg gtg ate ege age gag aac tte ace gac aac gec aag ace ate ate gtg eac etg aat gag age gtg eag ate (SEQ ID NO:11)

oligo 4 reverse (Mlu1): agt tgg gac gcg tgc agt tga tct gca cgc tct c (SEQ ID NO:12).

oligo 5 forward (Mlu1): gag agc gtg cag atc aac tgc acg cgt ccc (SEQ ID NO:13).

oligo 5: aac tgc acg cgt ccc aac tac aac aag cgc aag cgc atc cac atc ggc ccc ggg cgc gcc ttc tac acc acc aag aac atc atc ggc acc atc ctc cag gcc cac tgc aac atc tct aga (SEQ ID NO:14).

oligo 5 reverse: gtc gtt cca ctt ggc tct aga gat gtt gca (SEQ ID NO:15).

oligo 6 forward: gca aca tet eta gag eca agt gga acg ac (SEQ ID NO:16).

oligo 6: gcc aag tgg aac gac acc ctg cgc cag atc gtg agc aag ctg aag gag cag ttc aag aac aag acc atc gtg ttc ac cag agc agc ggc ggc gac ccc gag atc gtg atg cac agc ttc aac tgc ggc ggc (SEQ ID NO:17).

oligo 6 reverse (EcoR1): gca gta gaa gaa ttc gcc gcc gca gtt ga (SEQ ID NO:18).

oligo 7 forward (EcoR1): tca act gcg gcg gcg aat tct tct act gc (SEQ ID NO:19).

oligo 7: ggc gaa ttc ttc tac tgc aac acc agc ccc ctg ttc aac agc acc tgg aac ggc aac aac acc tgg aac aac acc acc ggc agc aac aac aat att acc ctc cag tgc aag atc aag cag atc aac atg tgg cag ggg ggg ggc aag gcc atg tac gcc ccc atc gag ggc cag atc cgg tgc agc agc (SEQ ID NO:20)

oligo 7 reverse: gca gac cgg tga tgt tgc tgc tgc acc gga tct ggc cct c (SEQ ID NO:21).

oligo 8 forward: cga ggg cca gat ccg gtg cag cag caa cat cac cgg tct g (SEQ ID NO:22).

oligo 8: aac atc acc ggt ctg ctg ctg acc cgc gac ggc ggc aag gac acc gac acc aac gac acc gaa atc ttc cgc ccc ggc ggc ggc gac atg cgc gac aac tgg aga tct gag ctg tac aag tac aag gtg gtg acg atc gag ccc ctg ggc gtg gcc ccc acc aag gcc aag cgc cgc gtg gtg cag cgc gag aag cgc (SEQ ID NO:23).

oligo 8 reverse (Not1): cgc ggg cgg ccg ctt tag cgc ttc tcg cgc tgc acc ac (SEQ ID NO:24).

The following oligonucleotides were used for the construction of the ratTHY-lenv gene.

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oligo 1 forward (BamH1/Hind3): cgc ggg gga tcc aag ctt acc atg att cca gta ata agt (SEQ ID NO:25).

oligo 1: atg aat cca gta ata agt ata aca tta tta tta agt gta tta caa atg agt aga gga caa aga gta ata agt tta aca gca tct tta gta aat caa aat ttg aga tta gat tgt aga cat gaa aat aat aca aat ttg cca ata caa cat gaa ttt tca tta acg (SEQ ID NO:26).

oligo 1 reverse (EcoR1/Mlu1): cgc ggg gaa ttc acg cgt taa tga aaa ttc atg ttg (SEQ ID NO:27).

oligo 2 forward (BamH1/Mlu1): cgc gga tcc acg cgt gaa aaa aaa cat (SEQ ID NO:28).

oligo 2: cgt gaa aaa aaa aaa cat gta tta agt gga aca tta gga gta cca gaa cat aca tat aga agt aga gta aat ttg ttt agt gat aga ttc ata aaa gta tta aca tat gca aat ttt aca aca aaa gat gaa gga gat tat atg tgt gag (SEQ ID NO:29).

oligo 2 reverse (EcoR1/Sac1): cgc gaa ttc gag ctc aca cat ata atc tcc (SEQ ID NO:30).

oligo 3 forward (BamH1/Sac1): cgc gga tcc gag ctc aga gta agt gga caa (SEQ ID NO:31).

oligo 3 reverse (EcoR1/Not1): cgc gaa ttc gcg gcc gct tca taa act tat aaa atc (SEQ ID NO:33).

Polymerase Chain Reaction

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Short, overlapping 15 to 25 mer oligonucleotides annealing at both ends were used to amplify the long oligonuclotides by polymerase chain reaction (PCR). Typical PCR conditions were: 35 cycles, 55°C annealing temperature, 0.2 sec extension time. PCR products were gel purified, phenol

extracted, and used in a subsequent PCR to generate longer fragments consisting of two adjacent small fragments. These longer fragments were cloned into a CDM7-derived plasmid containing a leader sequence of the CD5 surface molecule followed by a Nhe1/Pst1/Mlu1/EcoR1/BamH1 polylinker.

The following solutions were used in these reactions: 10x PCR buffer (500 mM KCl, 100 mM Tris HCl, pH 7.5, 8 mM MgCl₂, 2 mM each dNTP). The final buffer was complemented with 10% DMSO to increase fidelity of the Taq polymerase.

Small scale DNA preparation

Transformed bacteria were grown in 3 ml LB cultures for more than 10 6 hours or overnight. Approximately 1.5 ml of each culture was poured into 1.5 ml microfuge tubes, spun for 20 seconds to pellet cells and resuspended in 200 μ l of solution I. Subsequently 400 μ l of solution II and 300 μ l of solution III were added. The microfuge tubes were capped, mixed and spun for > 30 sec. Supernatants were transferred into fresh tubes and phenol extracted once. DNA 15 was precipitated by filling the tubes with isopropanol, mixing, and spinning in a microfuge for > 2 min. The pellets were rinsed in 70 % ethanol and resuspended in 50 μ l dH20 containing 10 μ l of RNAse A. The following media and solutions were used in these procedures: LB medium (1.0 % NaCl, 0.5% yeast extract, 1.0% trypton); solution I (10 mM EDTA pH 8.0); solution 20 II (0.2 M NaOH, 1.0% SDS); solution III (2.5 M KOAc, 2.5 M glacial aceatic acid); phenol (pH adjusted to 6.0, overlaid with TE); TE (10 mM Tris HCl, pH 7.5, 1 mM EDTA pH 8.0).

Large scale DNA preparation

One liter cultures of transformed bacteria were grown 24 to 36 hours (MC1061p3 transformed with pCDM derivatives) or 12 to 16 hours (MC1061 transformed with pUC derivatives) at 37°C in either M9 bacterial medium

(pCDM derivatives) or LB (pUC derivatives). Bacteria were spun down in 1 liter bottles using a Beckman J6 centrifuge at 4,200 rpm for 20 min. The pellet was resuspended in 40 ml of solution I. Subsequently, 80 ml of solution II and 40 ml of solution III were added and the bottles were shaken semivigorously until lumps of 2 to 3 mm size developed. The bottle was spun at 4,200 rpm for 5 min and the supernatant was poured through cheescoloth into a 250 ml bottle.

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Isopropanol was added to the top and the bottle was spun at 4,200 rpm for 10 min. The pellet was resuspended in 4.1 ml of solution I and added to 4.5 g of cesium chloride, 0.3 ml of 10 mg/ml ethidium bromide, and 0.1 ml of 1% Triton X100 solution. The tubes were spun in a Beckman J2 high speed centrifuge at 10,000 rpm for 5 min. The supernatant was transferred into Beckman Quick Seal ultracentrifuge tubes, which were then sealed and spun in a Beckman ultracentrifuge using a NVT90 fixed angle rotor at 80,000 rpm for > 2.5 hours. The band was extracted by visible light using a 1 ml syringe and 20 gauge needle. An equal volume of dH₂O was added to the extracted material. DNA was extracted once with n-butanol saturated with 1 M sodium chloride, followed by addition of an equal volume of 10 M ammonium acetate/ 1 mM EDTA. The material was poured into a 13 ml snap tube which was tehn filled to the top with absolute ethanol, mixed, and spun in a Beckman J2 centrifuge at 10,000 rpm for 10 min. The pellet was rinsed with 70% ethanol and resuspended in 0.5 to 1 ml of H₂O. The DNA concentration was determined by measuring the optical density at 260 nm in a dilution of 1:200 (1 $OD_{260} = 50$ μ g/ml).

The following media and buffers were used in these procedures: M9 bacterial medium (10 g M9 salts, 10 g casamino acids (hydrolyzed), 10 ml M9 additions, 7.5 μ g/ml tetracycline (500 μ l of a 15 mg/ml stock solution), 12.5 μ g/ml ampicillin (125 μ l of a 10 mg/ml stock solution); M9 additions (10 mM

CaCl₂, 100 mM MgSO₄, 200 μg/ml thiamine, 70% glycerol); LB medium (1.0 % NaCl, 0.5 % yeast extract, 1.0 % trypton); Solution I (10 mM EDTA pH 8.0); Solution II (0.2 M NaOH 1.0 % SDS); Solution III (2.5 M KOAc 2.5 M HOAc)

Sequencing

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Synthetic genes were sequenced by the Sanger dideoxynucleotide method. In brief, 20 to 50 μ g double-stranded plasmid DNA were denatured in 0.5 M NaOH for 5 min. Subsequently the DNA was precipitated with 1/10 volume of sodium acetate (pH 5.2) and 2 volumes of ethanol and centrifuged for 5 min. The pellet was washed with 70% ethanol and resuspended at a concentration of 1 μ g/ μ l. The annealing reaction was carried out with 4 μ g of template DNA and 40 ng of primer in 1x annealing buffer in a final volume of 10 μ l. The reaction was heated to 65°C and slowly cooled to 37°C.

In a separate tube 1 μ l of 0.1 M DTT, 2 μ l of labeling mix, 0.75 μ l of dH₂0, 1 μ l of [35 S] dATP (10 μ Ci), and 0.25 μ l of SequenaseTM (12 U/ μ l) were added for each reaction. Five μ l of this mix were added to each annealed primer-template tube and incubated for 5 min at room temperature. For each labeling reaction 2.5 μ l of each of the 4 termination mixes were added on a Terasaki plate and prewarmed at 37°C. At the end of the incubation period 3.5 μ l of labeling reaction were added to each of the 4 termination mixes. After 5 min, 4 μ l of stop solution were added to each reaction and the Terasaki plate was incubated at 80°C for 10 min in an oven. The sequencing reactions were run on 5% denaturing polyacrylamide gel. An acrylamide solution was prepared by adding 200 ml of 10x TBE buffer and 957 ml of dH₂0 to 100 g of acrylamide:bisacrylamide (29:1). 5% polyacrylamide 46% urea and 1x TBE gel was prepared by combining 38 ml of acrylamide solution and 28 g urea. Polymerization was initiated by the addition of 400 μ l of 10% ammonium

peroxodisulfate and 60 μ l of TEMED. Gels were poured using silanized glass plates and sharktooth combs and run in 1x TBE buffer at 60 to 100 W for 2 to 4 hours (depending on the region to be read). Gels were transferred to Whatman blotting paper, dried at 80 °C for about 1 hour, and exposed to x-ray film at room temperature. Typically exposure time was 12 hours. The following solutions were used in these procedures: 5x Annealing buffer (200 mM Tris HCl, pH 7.5, 100 mM MgCl₂, 250 mM NaCl); Labelling Mix (7.5 μ M each dCTP, dGTP, and dTTP); Termination Mixes (80 μ M each dNTP, 50 mM NaCl, 8 μ M ddNTP (one each)); Stop solution (95% formamide, 20 mM EDTA, 0.05 % bromphenol blue, 0.05 % xylencyanol); 5x TBE (0.9 M Tris borate, 20 mM EDTA); Polyacrylamide solution (96.7 g polyacrylamide, 3.3 g bisacrylamide, 200 ml 1x TBE, 957 ml dH₂O).

RNA isolation

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Cytoplasmic RNA was isolated from calcium phosphate transfected 293T cells 36 hours post transfection and from vaccinia infected Hela cells 16 hours post infection essentially as described by Gilman. (Gilman Preparation of cytoplasmic RNA from tissue culture cells. In Current Protocols in Molecular Biology, Ausubel et al., eds., Wiley & Sons, New York, 1992). Briefly, cells were lysed in 400 μ l lysis buffer, nuclei were spun out, and SDS and proteinase K were added to 0.2% and 0.2 mg/ml respectively. The cytoplasmic extracts were incubated at 37°C for 20 min, phenol/chloroform extracted twice, and precipitated. The RNA was dissolved in 100 μ l buffer I and incubated at 37°C for 20 min. The reaction was stopped by adding 25 μ l stop buffer and precipitated again.

The following solutions were used in this procedure: Lysis Buffer (TRUSTEE containing with 50 mM Tris pH 8.0, 100 mM NaCl, 5 mM MgCl₂, 0.5% NP40); Buffer I (TRUSTEE buffer with 10 mM MgCl₂, 1 mM DTT, 0.5

U/µl placental RNAse inhibitor, 0.1 U/µl RNAse free DNAse I); Stop buffer (50 mM EDTA 1.5 M NaOAc 1.0% SDS).

Slot blot analysis

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For slot blot analysis 10 μ g of cytoplasmic RNA was dissolved in 50 μ l dH₂O to which 150 μ l of 10x SSC/18% formaldehyde were added. The solubilized RNA was then incubated at 65°C for 15 min and spotted onto with a slot blot apparatus. Radioactively labeled probes of 1.5 kb gp120IIIb and syngp120mn fragments were used for hybridization. Each of the two fragments was random labeled in a 50 μ l reaction with 10 μ l of 5x oligo-labeling buffer, 8 μ l of 2.5 mg/ml BSA, 4 μ l of [\propto ³²P]-dCTP (20 uCi/ μ l; 6000 Ci/mmol), and 5 U of Klenow fragment. After 1 to 3 hours incubation at 37°C 100 μ l of TRUSTEE were added and unincorporated [x32P]-dCTP was eliminated using G50 spin column. Activity was measured in a Beckman beta-counter, and equal specific activities were used for hybridization. Membranes were prehybridized for 2 hours and hybridized for 12 to 24 hours at 42°C with 0.5 x 106 cpm probe per ml hybridization fluid. The membrane was washed twice (5 min) with washing buffer I at room temperature, for one hour in washing buffer Il at 65°C, and then exposed to x-ray film. Similar results were obtained using a 1.1 kb Not1/Sfi1 fragment of pCDM7 containing the 3 untranslated region. Control hybridizations were done in parallel with a random-labeled human beta-actin probe. RNA expression was quantitated by scanning the hybridized nitrocellulose membranes with a Magnetic Dynamics phosphorimager.

The following solutions were used in this procedure:

5x Oligo-labeling buffer (250 mM Tris HCl, pH 8.0, 25 mM MgCl₂, 5 mM β-mercaptoethanol, 2 mM dATP, 2 mM dGTP, mM dTTP, 1 M Hepes pH 6.6, 1 mg/ml hexanucleotides [dNTP]6); Hybridization Solution (.05 M sodium phosphate, 250 mM NaCl, 7% SDS, 1 mM EDTA, 5% dextrane sulfate, 50%

formamide, 100 μ g/ml denatured salmon sperm DNA); Washing buffer I (2x SSC,

0.1% SDS); Washing buffer II (0.5x SSC, 0.1 % SDS); 20x SSC (3 M NaCl, 0.3 M Na₃citrate, pH adjusted to 7.0).

Vaccinia recombination

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Vaccinia recombination used a modification of the of the method described by Romeo and Seed (Romeo and Seed, Cell, 64: 1037, 1991). Briefly, CV1 cells at 70 to 90% confluency were infected with 1 to 3 μ 1 of a wild-type vaccinia stock WR (2 x 108 pfu/ml) for 1 hour in culture medium without calf serum. After 24 hours, the cells were transfected by calcium phosphate with 25 µg TKG plasmid DNA per dish. After an additional 24 to 48 hours the cells were scraped off the plate, spun down, and resuspended in a volume of 1 ml. After 3 freeze/thaw cycles trypsin was added to 0.05 mg/ml and lysates were incubated for 20 min. A dilution series of 10, 1 and 0.1 μ l of this lysate was used to infect small dishes (6 cm) of CV1 cells, that had been pretreated with 12.5 µg/ml mycophenolic acid, 0.25 mg/ml xanthin and 1.36 mg/ml hypoxanthine for 6 hours. Infected cells were cultured for 2 to 3 days, and subsequently stained with the monoclonal antibody NEA9301 against gp120 and an alkaline phosphatase conjugated secondary antibody. Cells were incubated with 0.33 mg/ml NBT and 0.16 mg/ml BCIP in AP-buffer and finally overlaid with 1% agarose in PBS. Positive plaques were picked and resuspended in 100 μ l Tris pH 9.0. The plaque purification was repeated once. To produce high titer stocks the infection was slowly scaled up. Finally, one large plate of Hela cells was infected with half of the virus of the previous round. Infected cells were detached in 3 ml of PBS, lysed with a Dounce homogenizer and cleared from larger debris by centrifugation. VPE-8 recombinant vaccinia stocks were kindly provided by the AIDS repository,

Rockville, MD, and express HIV-1 IIIB gp120 under the 7.5 mixed early/late promoter (Earl et al., <u>J. Virol.</u>, 65:31, 1991). In all experiments with recombinant vaccina cells were infected at a multiplicity of infection of at least 10.

The following solution was used in this procedure:

AP buffer (100 mM Tris HCl, pH 9.5, 100 mM NaCl, 5 mM MgCl₂)

Cell culture

The monkey kidney carcinoma cell lines CV1 and Cos7, the human kidney carcinoma cell line 293T, and the human cervix carcinoma cell line Hela were obtained from the American Tissue Typing Collection and were maintained in supplemented IMDM. They were kept on 10 cm tissue culture plates and typically split 1:5 to 1:20 every 3 to 4 days. The following medium was used in this procedure:

Supplemented IMDM (90% Iscove's modified Dulbecco Medium, 10% calf serum, iron-complemented, heat inactivated 30 min 56°C, 0.3 mg/ml L-glutamine, 25 μ g/ml gentamycin 0.5 mM β -mercaptoethanol (pH adjusted with 5 M NaOH, 0.5 ml)).

Transfection

Calcium phosphate transfection of 293T cells was performed by

slowly adding and under vortexing 10 μg plasmid DNA in 250 μl 0.25 M

CaCl₂ to the same volume of 2x HEBS buffer while vortexing. After incubation for 10 to 30 min at room temperature the DNA precipitate was added to a small dish of 50 to 70% confluent cells. In cotransfection experiments with rev, cells were transfected with 10 μg gp120IIIb,

gp120IIIbrre, syngp120mnrre or rTHY-lenveg1rre and 10 μg of pCMVrev or CDM7 plasmid DNA.

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The following solutions were used in this procedure: 2x HEBS buffer (280 mM NaCl, 10 mM KCl, 1.5 mM sterile filtered); 0.25 mM CaCl₂ (autoclaved).

Immunoprecipitation

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After 48 to 60 hours medium was exchanged and cells were incubated for additional 12 hours in Cys/Met-free medium containing 200 µCi of ³⁵S-translabel. Supernatants were harvested and spun for 15 min at 3000 rpm to remove debris. After addition of protease inhibitors leupeptin, aprotinin and PMSF to 2.5 μg/ml, 50 μg/ml, 100 μg/ml respectively, 1 ml of supernatant was incubated with either 10 µl of packed protein A sepharose alone (rTHYlenveglrre) or with protein A sepharose and 3 µg of a purified CD4/immunoglobulin fusion protein (kindly provided by Behring) (all gp120 constructs) at 4°C for 12 hours on a rotator. Subsequently the protein A beads were washed 5 times for 5 to 15 min each time. After the final wash 10 µl of loading buffer containing was added, samples were boiled for 3 min and applied on 7% (all gp120 constructs) or 10% (rTHY-lenveg1rre) SDS polyacrylamide gels (TRIS pH 8.8 buffer in the resolving, TRIS pH 6.8 buffer in the stacking gel, TRIS-glycin running buffer, Maniatis et al., supra 1989). Gels were fixed in 10% acetic acid and 10 % methanol, incubated with Amplify for 20 min, dried and exposed for 12 hours.

The following buffers and solutions were used in this procedure:

Wash buffer (100 mM Tris, pH 7.5, 150 mM NaCl, 5 mM CaCl₂, 1% NP-40);

5x Running Buffer (125 mM Tris, 1.25 M Glycin, 0.5% SDS); Loading buffer (10 % glycerol, 4% SDS, 4% β-mercaptoethanol, 0.02 % bromphenol blue).

Immunofluorescence

293T cells were transfected by calcium phosphate coprecipitation and analyzed for surface THY-1 expression after 3 days. After detachment

with 1 mM EDTA/PBS, cells were stained with the monoclonal antibody OX-7 in a dilution of 1:250 at 4°C for 20 min, washed with PBS and subsequently incubated with a 1:500 dilution of a FITC-conjugated goat anti-mouse immunoglobulin antiserum. Cells were washed again, resuspended in 0.5 ml of a fixing solution, and analyzed on a EPICS XL cytofluorometer (Coulter).

The following solutions were used in this procedure:
PBS (137 mM NaCl, 2.7 mM KCl, 4.3 mM Na₂HPO₄, 1.4 mM KH₂PO₄, pH adjusted to 7.4); Fixing solution (2% formaldehyde in PBS).

ELISA

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The concentration of gp120 in culture supernatants was determined using CD4-coated ELISA plates and goat anti-gp120 antisera in the soluble phase. Supernatants of 293T cells transfected by calcium phosphate were harvested after 4 days, spun at 3000 rpm for 10 min to remove debris and incubated for 12 hours at 4°C on the plates. After 6 washes with PBS 100 μl of goat anti-gp120 antisera diluted 1:200 were added for 2 hours. The plates were washed again and incubated for 2 hours with a peroxidase-conjugated rabbit anti-goat IgG antiserum 1:1000. Subsequently the plates were washed and incubated for 30 min with 100 μl of substrate solution containing 2 mg/ml ophenylenediamine in sodium citrate buffer. The reaction was finally stopped with 100 μl of 4 M sulfuric acid. Plates were read at 490 nm with a Coulter microplate reader. Purified recombinant gp120IIIb was used as a control. The following buffers and solutions were used in this procedure: Wash buffer (0.1% NP40 in PBS); Substrate solution (2 mg/ml o-phenylenediamine in sodium citrate buffer).

EXAMPLE 2

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A Synthetic Green Fluorescent Protein Gene

The efficacy of codon replacement for gp120 suggests that replacing non-preferred codons with less preferred codons or preferred codons (and replacing less preferred codons with preferred codons) will increase expression in mammalian cells of other proteins, e.g., other eukaryotic proteins.

The green fluorescent protein (GFP) of the jellyfish Aequorea victoria (Ward, Photochem. Photobiol. 4:1, 1979; Prasher et al., Gene 111:229, 1992; Cody et al., Biochem. 32:1212, 1993) has attracted attention recently for its possible utility as a marker or reporter for transfection and lineage studies (Chalfie et al., Science 263:802, 1994).

Examination of a codon usage table constructed from the native coding sequence of GFP showed that the GFP codons favored either A or U in the third position. The bias in this case favors A less than does the bias of gp120, but is substantial. A synthetic gene was created in which the natural GFP sequence was re-engineered in much the same manner as for gp120 (FIG. 11; SEQ ID NO:40). In addition, the translation initiation sequence of GFP was replaced with sequences corresponding to the translational initiation consensus. The expression of the resulting protein was contrasted with that of the wild type sequence, similarly engineered to bear an optimized translational initiation consensus (FIG. 10B and FIG. 10C). In addition, the effect of inclusion of the mutation Ser 65-Thr, reported to improve excitation efficiency of GFP at 490 nm and hence preferred for fluorescence microscopy (Heim et al., Nature 373:663, 1995), was examined (FIG. 10D). Codon engineering conferred a significant increase in expression efficiency (an concomitant percentage of cells apparently positive for transfection), and the combination of

the Ser 65-Thr mutation and codon optimization resulted in a DNA segment encoding a highly visible mammalian marker protein (FIG. 10D).

The above-described synthetic green fluorescent protein coding sequence was assembled in a similar manner as for gp120 from six fragments of approximately 120 bp each, using a strategy for assembly that relied on the ability of the restriction enzymes BsaI and BbsI to cleave outside of their recognition sequence. Long oligonucleotides were synthesized which contained portions of the coding sequence for GFP embedded in flanking sequences encoding EcoRI and BsaI at one end, and BamHI and BbsI at the other end. Thus, each oligonucleotide has the configuration EcoRI/BsaI/GFP fragment/BbsI/BamHI. The restriction site ends generated by the BsaI and BbsI sites were designed to yield compatible ends that could be used to join adjacent GFP fragments. Each of the compatible ends were designed to be unique and non-selfcomplementary. The crude synthetic DNA segments were amplified by PCR, inserted between EcoRI and BamHI in pUC9, and sequenced. Subsequently the intact coding sequence was assembled in a six fragment ligation, using insert fragments prepared with BsaI and BbsI. Two of six plasmids resulting from the ligation bore an insert of correct size, and one contained the desired full length sequence. Mutation of Ser65 to Thr was accomplished by standard PCR based mutagenesis, using a primer that overlapped a unique BssSI site in the synthetic GFP.

Codon optimization as a strategy for improved expression in mammalian cells

The data presented here suggest that coding sequence re-engineering may have general utility for the improvement of expression of mammalian and non-mammalian eukaryotic genes in mammalian cells. The results obtained here with three unrelated proteins: HIV gp120, the rat cell surface antigen Thy-

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1 and green fluorescent protein from Aequorea victoria, and human Factor VIII (see below) suggest that codon optimization may prove to be a fruitful strategy for improving the expression in mammalian cells of a wide variety of eukaryotic genes.

5 **EXAMPLE III**

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Design of a Codon-Optimized Gene Expressing Human Factor VIII Lacking the Central B Domain

A synthetic gene was designed that encodes mature human Factor VIII lacking amino acid residues 760 to 1639, inclusive (residues 779 to 1658, inclusive, of the precursor). The synthetic gene was created by choosing codons corresponding to those favored by highly expressed human genes. Some deviation from strict adherence to the favored residue pattern was made to allow unique restriction enzyme cleavage sites to be introduced throughout the gene to facilitate future manipulations. For preparation of the synthetic gene the sequence was then divided into 28 segments of 150 basepairs, and a 29th segment of 161 basepairs.

The a synthetic gene expressing human Factor VIII lacking the central B domain was constructed as follows. Twenty-nine pairs of template oligonucleotides (see below) were synthesized. The 5' template oligos were 105 bases long and the 3' oligos were 104 bases long (except for the last 3' oligo, which was 125 residues long). The template oligos were designed so that each annealing pair composed of one 5' oligo and one 3' oligo, created a 19 basepair double-stranded regions.

To facilitate the PCR and subsequent manipulations, the 5' ends of
the oligo pairs were designed to be invariant over the first 18 residues, allowing
a common pair of PCR primers to be used for amplification, and allowing the
same PCR conditions to be used for all pairs. The first 18 residues of each 5'

member of the template pair were cgc gaa ttc gga aga ccc (SEQ ID NO:110) and the first 18 residues of each 3' member of the template pair were: ggg gat cct cac gtc tca (SEQ ID NO:43).

Pairs of oligos were annealed and then extended and amplified by PCR in a reaction mixture as follows: templates were annealed at 200 μg/ml each in PCR buffer (10 mM Tris-HCl, 1.5 mM MgCl₂, 50 mM KCl, 100 μg/ml gelatin, pH 8.3). The PCR reactions contained 2 ng of the annealed template oligos, 0.5 μg of each of the two 18-mer primers (described below), 200 μM of each of the deoxynucleoside triphosphates, 10% by volume of DMSO and PCR buffer as supplied by Boehringer Mannheim Biochemicals, in a final volume of 50 μl. After the addition of Taq polymerase (2.5 units, 0.5 μl; Boehringer Mannheim Biochemicals) amplifications were conducted on a Perkin-Elmer Thermal Cycler for 25 cycles (94°C for 30 sec, 55°C for 30 sec, and 72°C for 30 sec). The final cycle was followed by a 10 minute extension at 72°C.

The amplified fragments were digested with EcoRI and BamHI (cleaving at the 5' and 3' ends of the fragments respectively) and ligated to a pUC9 derivative cut with EcoRI and BamHI.

Individual clones were sequenced and a collection of plasmids corresponding to the entire desired sequence was identified. The clones were then assembled by multifragment ligation taking advantage of restriction sites at the 3' ends of the PCR primers, immediately adjacent to the amplified sequence. The 5' PCR primer contained a BbsI site, and the 3' PCR primer contained a BsmBI site, positioned so that cleavage by the respective enzymes preceded the first nucleotide of the amplified portion and left a 4 base 5' overhang created by the first 4 bases of the amplified portion. Simultaneous digestion with BbsI and BsmBI thus liberated the amplified portion with unique 4 base 5' overhangs at each end which contained none of the primer sequences.

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In general these overhangs were not self-complementary, allowing multifragment ligation reactions to produce the desired product with high efficiency. The unique portion of the first 28 amplified oligonucleotide pairs was thereby 154 basepairs, and after digestion each gave rise to a 150 bp fragment with unique ends. The first and last fragments were not manipulated in this manner, however, since they had other restriction sites designed into them to facilitate insertion of the assembled sequence into an appropriate mammalian expression vector. The actual assembly process proceded as follows.

10 Assembly of the Synthetic Factor VIII Gene

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Step 1: 29 Fragments Assembled to Form 10 Fragments.

The 29 pairs of oligonucleotides, which formed segments 1 to 29 when base-paired, are described below.

Plasmids carrying segments 1, 5, 9, 12, 16, 20, 24 and 27 were digested with EcoR1 and BsmBI and the 170 bp fragments were isolated; plasmids bearing segments 2, 3, 6, 7, 10, 13, 17, 18, 21, 25, and 28 were digested with BbsI and BsmBI and the 170 bp fragments were isolated; and plasmids bearing segments 4, 8, 11, 14, 19, 22, 26 and 29 were digested with EcoR1 and BbsI and the 2440 bp vector fragment was isolated. Fragments bearing segments 1, 2, 3 and 4 were then ligated to generate segment "A"; fragments bearing segments 5, 6, 7 and 8 were ligated to generate segment "B"; fragments bearing segments 12, 13, and 11 were ligated to generate segment "C"; fragments bearing segments 16, 17, 18 and 19 were ligated to generate segment "D"; fragments bearing segments 20, 21 and 22 were ligated to generate segment "G"; fragments bearing segments 24, 25 and 26 were ligated

to generate segment "I"; and fragments bearing segments 27, 28 and 29 were ligated to generate segment "J".

Step 2: Assembly of the 10 resulting Fragments from Step 1 to Three Fragments.

Plasmids carrying the segments "A", "D" and "G" were digested with EcoRI and BsmBI, plasmids carrying the segments B, 15, 23, and I were digested with BbsI and BsmBI, and plasmids carrying the segments C, F, and J were digested with EcoRI and BbsI. Fragments bearing segments A, B, and C were ligated to generate segment "K"; fragments bearing segments D, 15, and F were ligated to generate segment "O"; and fragments bearing segments G, 23, I, and J were ligated to generate segment "P".

Step 3: Assembly of the Final Three Pieces.

The plasmid bearing segment K was digested with EcoRI and BsmBI, the plasmid bearing segment O was digested with BbsI and BsmBI, and the plasid bearing segment P was digested with EcoRI and BbsI. The three resulting fragments were ligated to generate segments.

Step 4: Insertion of the Synthetic Gene in a Mammalian Expression Vector.

The plasmid bearing segment S was digested with Nhel and Notl and inserted between Nhel and Eagl sites of plasmid CD5lNEg1 to generate plasmid cd5lsf8b.

Sequencing and Correction of the Synthetic Factor VIII Gene

After assembly of the synthetic gene it was discovered that there were two undesired residues encoded in the sequence. One was an Arg residue at 749, which is present in the GenBank sequence entry originating from Genentech but is not in the sequence reported by Genentech in the literature. The other was an Ala residue at 146, which should have been Pro. This

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mutation arose at an unidentified step subsequent to the sequencing of the 29 constituent fragments. The Pro749Arg mutation was corrected by incorporating the desired change in a PCR primer (ctg ctt ctg acg cgt gct ggg gtg gcg gga gtt; SEQ ID NO:44) that included the MluI site at position 2335 of the sequence below (sequence of HindIII to NotI segment) and amplifying between that primer and a primer (ctg ctg aaa gtc tcc agc tgc; SEQ ID NO:44) 5' to the SgrAI site at 2225. The SgrAI to MluI fragment was then inserted into the expression vector at the cognate sites in the vector, and the resulting correct sequence change verified by sequencing. The Pro146Ala mutation was corrected by incorporating the desired sequence change in an oligonucleotide (ggc agg tgc tta agg aga acg gcc cta tgg cca; SEQ ID NO:46) bearing the AfIII site at residue 504, and amplifying the fragment resulting from PCR reaction between that oligo and the primer having sequence cgt tgt tct tca tac gcg tct ggg gct cct cgg ggc (SEQ ID NO:109), cutting the resulting PCR fragment with AfIII and AvrII at (residue 989), inserting the corrected fragment into the expression vector and confirming the construction by sequencing. Construction of a Matched Native Gene Expressing Human Factor VIII Lacking the Central B Domain

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20 the native codon sequence was constructed by introducing NheI at the 5' end of the mature coding sequence using primer cgc caa ggg cta gcc gcc acc aga aga tac tac ctg ggt (SEQ ID NO:47), amplifying between that primer and the primer att cgt agt tgg ggt tcc tct gga cag (corresponding to residues 1067 to 1093 of the sequence shown below), cutting with NheI and AfIII (residue 345 in the sequence shown below) and inserting the resulting fragment into an appropriately cleaved plasmid bearing native Factor VIII. The B domain deletion was created by overlap PCR using ctg tat ttg atg aga acc g,

(corresponding to residues 1813 to 1831 below) and caa gac tgg tgg ggt ggc att aaa ttg ctt t (SEQ ID NO:48) (2342 to 2372 on complement below) for the 5' end of the overlap, and aat gcc acc cca cca gtc ttg aaa cgc ca (SEQ ID NO:49) (2352 to 2380 on sequence below) and cat ctg gat att gca ggg ag (SEQ ID NO:50) (3145 to 3164). The products of the two individual PCR reactions were then mixed and reamplified by use of the outermost primers, the resulting fragment cleaved by Asp718 (KpnI isoschizomer, 1837 on sequence below) and PflMI (3100 on sequence below), and inserted into the appropriately cleaved expression plasmid bearing native Factor VIII.

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The complete sequence (SEQ ID NO:41) of the native human factor VIII gene deleted for the central B region is presented in Figure 12. The complete sequence (SEQ ID NO:42) of the synthetic Factor VIII gene deleted for the central B region is presented in Figure 13.

Preparation and assay of expression plasmids

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Two independent plasmid isolates of the native, and four independent isolates of the synthetic Factor VIII expression plasmid were separately propagated in bacteria and their DNA prepared by CsCl buoyant density centrifugation followed by phenol extraction. Analysis of the supernatants of COS cells transfected with the plasmids showed that the synthetic gene gave rise to approximately four times as much Factor VIII as did the native gene.

COS cells were then transfected with 5 µg of each factor VIII construct per 6 cm dish using the DEAE-dextran method. At 72 hours post-transfection, 4 ml of fresh medium containing 10% calf serum was added to each plated. A sample of media was taken from each plate 12 hr later. Samples were tested by ELISA using mouse anti-human factor VIII light chain monoclonal antibody and peroxidase-conjugated goat anti-human factor VIII

polyclonal antibody. Purified human plasma factor VIII was used as a standard. Cells transfected with the synthetic Factor V111 gene construct expressed 138 ± 20.2 ng/ml (equivalent ng/ml non-deleted Factor VIII) of Factor VIII (n=4) while the cells transfected with the native Factor VIII gene expressed 33.5 ± 0.7 ng/ml (equivalent ng/ml non-deleted Factor VIII) of Factor VIII (n=2).

The following template oligonucleotides were used for construction of the synthetic Factor VIII gene.

r1 bbs 1 for (gcta)

cgc gaa ttc gga aga ccc gct agc cgc cac l rl
ccg ccg cta cta cct ggg cgc cgt gga gct
gtc ctg gga cta cat gca gag cga cct ggg
cga gct ccc cgt gga (SEQ ID NO:51)

ggg gat cet cae gte tea ggt ttt ett gta 1 bam

15 cae cae get ggt gtt gaa ggg gaa get ett

ggg cae geg ggg ggg gaa geg gge gte cae

ggg gag ete gee ca (SEQ ID NO:52)

rl bbs 2 for (aacc)

cgc gaa ttc gga aga ccc aac cct gtt cgt

20 gga gtt cac cga cca cct gtt caa cat tgc
caa gcc gcg ccc ccc ctg gat ggg cct gct
ggg ccc cac cat cca (SEQ ID NO:53)

2 r1

ggg gat cct cac gtc tca gtg cag gct gac ggg gtg gct ggc cat gtt ctt cag ggt gat cac cac ggt gtc gta cac ctc ggc ctg gat ggt ggg gcc cag ca (SEQ ID NO:54) 2 bam

r1 bbs 3 for (gcac)

cgc gaa ttc gga aga ccc gca cgc cgt ggg cgt gag cta ctg gaa ggc cag cga ggg cgc cga gta cga cga cca gac gtc cca gcg cga gaa gga gga cga caa (SEQ ID NO:55) 3 r1

ggg gat cct cac gtc tca gct ggc cat agg
gcc gtt ctc ctt aag cac ctg cca cac gta
ggt gtg gct ccc ccc cgg gaa cac ctt gtc
gtc ctc ctt ctc gc (SEQ ID NO:56)

3 bam

rl bbs 4 for (cagc)

cgc gaa ttc gga aga ccc cag cga ccc cct
gtg cct gac cta cag cta cct gag cca cgt
gga cct ggt gaa gga tct gaa cag cgg gct
gat cgg cgc cct gct (SEQ ID NO:57)

4 r1

ggg gat cet cae gte tea gaa eag eag gat

gaa ett gtg eag ggt etg ggt ttt ete ett

gge eag get gee ete geg aca eac eag eag

gge gee gat eag ee (SEQ ID NO:58)

4 bam

5

rl bbs 5 for (gttc)

cgc gaa ttc gga aga ccc gtt cgc cgt gtt

cga cga ggg gaa gag ctg gca cag cga gac

taa gaa cag cct gat gca gga ccg cga cgc
cgc cag cgc ccg cgc (SEQ ID NO:59)

ggg gat cet cae gte tea gtg gea gee gat cag gee ggg cag get geg gtt cae gta gee gtt aac ggt gtg cat ett ggg eea gge geg gge get gge gge gt (SEQ ID NO:60) 5 bam

rl bbs 6 for (ccac)

cgc gaa ttc gga aga ccc cca ccg caa gag
cgt gta ctg gca cgt cat cgg cat ggg cac
cac ccc tga ggt gca cag cat ctt cct gga
ggg cca cac ctt cct (SEQ ID NO:61)

6 r1

agt cag gat cct cac gtc tca cag ggt ctg ggc
agt cag gaa ggt gat ggg gct gat ctc cag
gct ggc ctg gcg gtg gtt gcg cac cag gaa
ggt gtg gcc ctc ca (SEQ ID NO:62)

6 bam

rl bbs 7 for (cctg)

cct agg cca gtt cct gct gtt ctg cca cat
cag cag cca gca cga cga cgg cat gga ggc
tta cgt gaa ggt gga (SEQ ID NO:63)

7 rl

ggg gat cet cae gte tea gte gte gta gte ete gge ete ete gtt gtt ett eat geg cag etg ggg ete ete ggg gea get gte eae ett eae gta age et (SEQ ID NO:64)

7 bam

5

rl bbs 8 for (cgac)

cgc gaa ttc gga aga ccc cga cct gac cga cag cga gat gga tgt cgt acg ctt cga cga cga caa cag ccc cag ctt cat cca gat ccg cag cgt ggc caa gaa (SEQ ID NO:65)

8 rl

ggg gat cet cae gte tea tae tag egg gge
gta gte cea gte ete ete ete gge gge gat
gta gtg cae cea ggt ett agg gtg ett ett
gge cae get geg ga (SEQ ID NO:66)

8 bam

r1 bbs 9 for (agta)

cgc gaa ttc gga aga ccc agt act ggc ccc
cga cga ccg cag cta caa gag cca gta cct
gaa caa cgg ccc cca gcg cat cgg ccg caa
gta caa gaa ggt gcg (SEQ ID NO:67)

9 r1

ggg gat cet cae gte tea gag gat gee gga

cte gtg etg gat gge ete geg ggt ett gaa

agt ete gte ggt gta gge eat gaa geg eae

ett ett gta ett ge (SEQ ID NO:68)

9 bam

5

rl bbs 10 for (cctc)

cgc gaa ttc gga aga ccc cct cgg ccc cct gct gta cgg cga ggt ggg cga cac cct gct gat cat ctt caa gaa cca ggc cag cag gcc cta caa cat cta ccc (SEQ ID NO:69)

ggg gat cet cae gte tea ett cag gtg ett
cae gee ett ggg cag geg geg get gta cag
ggg geg cae gte ggt gat gee gtg ggg gta
gat gtt gta ggg ce (SEQ ID NO:70)

10 ham

10 rl

rl bbs 11 for (gaag)

cgc gaa ttc gga aga ccc gaa gga ctt ccc cat cct gcc cgg cga gat ctt caa gta caa gtg gac cgt gac cgt gga gga cgg ccc cac caa gag cga ccc ccg (SEQ ID NO:71)

11 rl

ggg gat cet cae gte tea gee gat eag tee
gga gge eag gte geg ete eat gtt eae gaa
get get gta gta geg ggt eag gea geg ggg
gte get ett ggt gg (SEQ ID NO:72)

11 bam

12 r1

rl bbs 12 for (cggc)

cgc gaa ttc gga aga ccc cgg ccc cct gct
gat ctg cta caa gga gag cgt gga cca gcg
cgg caa cca gat cat gag cga caa gcg caa
cgt gat cct gtt cag (SEQ ID NO:73)

ggg gat cct cac gtc tca agc ggg gtt ggg cag gaa gcg ctg gat gtt ctc ggt cag ata cca gct gcg gtt ctc gtc gaa cac gct gaa cag gat cac gtt gc (SEQ ID NO:74) 12 bam

r1 bbs 13 for (cgct)

cgc gaa ttc gga aga ccc cgc tgg cgt gca gct gga aga tcc cga gtt cca ggc cag caa cat cat gca cag cat caa cgg cta cgt gtt cga cag cct gca gct (SEQ ID NO:75) 13 rl

ctg ggc gcc gat gct cag gat gta cca gta
ggc cac ctc atg cag gca cac gct cag ctg
cag gct gtc gaa ca (SEQ ID NO:76)

13 bam

rl bbs 14 for (cctg)

cgc gaa ttc gga aga ccc cct gag cgt gtt
ctt ctc cgg gta tac ctt caa gca caa gat
ggt gta cga gga cac cct gac cct gtt ccc
ctt ctc cgg cga gac (SEQ ID NO:77)

14 r1

ggg gat cet cae gte tea gtt geg gaa gte

get gtt gtg gea gee eag aat eea eag gee
ggg gtt ete eat aga eat gaa eae agt ete
gee gga gaa ggg ga (SEQ ID NO:78)

14 bam

5

rl bbs 15 for (caac)

cgc gaa ttc gga aga ccc caa ccg cgg cat gac tgc cct gct gaa agt ctc cag ctg cga caa gaa cac cgg cga cta cta cga gga cag cta cga gga cat ctc (SEQ ID NO:79)

15 rl

ggg gat cet cae gte tea geg gtg geg gga gtt ttg gga gaa gga geg ggg ete gat gge gtt gtt ett gga eag eag gta gge gga gat gte ete gta get gt (SEQ ID NO:80)

15 bam

rl bbs 16 for (ccgc)

cgc gaa ttc gga aga ccc ccg cag cac gcg tca gaa gca gtt caa cgc cac ccc ccc cgt gct gaa gcg cca cca gcg cga gat cac ccg cac cac cct gca aag (SEQ ID NO:81) 16 r1

ggg gat cet cae gte tea gat gte gaa gte
ete ett ett eat ete eae get gat ggt gte
gte gta gte gat ete ete etg gte get ttg
eag ggt ggt geg gg (SEQ ID NO:82)

16 bam

rl bbs 17 for (catc)

20 cgc gaa ttc gga aga ccc cat cta cga cga gga cga gaa cca gag ccc ccg ctc ctt cca aaa gaa aac ccg cca cta ctt cat cgc cgc cgt gga gcg cct gtg (SEQ ID NO:83)

17 r1

ggg gat cet cae gte tea etg ggg eae get gee get etg gge geg gtt geg eag gae gtg ggg get get get eat gee gta gte cea eag geg etc eae gge gg (SEQ ID NO:84) 17 bam

5 r1 bbs 18 for (ccag)

cgc gaa ttc gga aga ccc cca gtt caa gaa ggt ggt gtt cca gga gtt cac cga cgg cag ctt cac cca gcc cct gta ccg cgg cga gct gaa cga gca cct ggg (SEQ ID NO:85) 18 rl

ggg gat cet cae gte tea gge ttg gtt geg
gaa ggt cae cat gat gtt gte ete eae ete
gge geg gat gta ggg gee gag eag gee eag
gtg ete gtt eag et (SEQ ID NO:86)

18 bam

rl bbs 19 for (agcc)

cta ctc ctt cta ctc ctc cct gat cag ctc ccg gcc
cta ctc ctt cta ctc ctc cct gat cag cta
cga gga gga cca gcg cca ggg cgc cga gcc
ccg caa gaa ctt cgt (SEQ ID NO:87)

19 rl

ggg gat cet cae gte tea ete gte ett ggt

20 ggg gge eat gtg gtg etg eae ett eea gaa
gta ggt ett agt ete gtt ggg ett eae gaa
gtt ett geg ggg et (SEQ ID NO:88)

19 bam

5

rl bbs 20 for (cgag)

cgc gaa ttc gga aga ccc cga gtt cga ctg
caa ggc ctg ggc cta ctt cag cga cgt gga
cct gga gaa gga cgt gca cag cgg cct gat
cgg ccc cct gct ggt (SEQ ID NO:89)

20 r1

ggg gat cet cae gte tea gaa cag gge aaa tte etg cae agt cae etg eet eee gtg ggg ggg gtt cag ggt gtt ggt gtg gea cae cag cag ggg gee gat ca (SEQ ID NO:90) 20 bam

rl bbs 21 for (gttc)

cgc gaa ttc gga aga ccc gtt ctt cac cat ctt cga cga gac taa gag ctg gta ctt cac cga gaa cat gga gcg caa ctg ccg cgc ccc ctg caa cat cca gat (SEQ ID NO:91) 21 r1

ggg gat cet cae gte tea eag ggt gte eat
gat gta gee gtt gat gge gtg gaa geg gta
gtt ete ett gaa ggt ggg ate tte eat etg
gat gtt gea ggg gg (SEQ ID NO:92)

21 bam

rl bbs 22 for (cctg)

20 cgc gaa ttc gga aga ccc cct gcc cgg cct
ggt gat ggc cca gga cca gcg cat ccg ctg
gta cct gct gtc tat ggg cag caa cga gaa
cat cca cag cat cca (SEQ ID NO:93)

22 rl

ggg gat cct cac gtc tca gta cag gtt gta cag ggc cat ctt gta ctc ctc ctt ctt gcg cac ggt gaa aac gtg gcc gct gaa gtg gat gct gtg gat gtt ct (SEQ ID NO:94)

22 bam

5

rl bbs 23 for (gtac)

cgc gaa ttc gga aga ccc gta ccc cgg cgt gtt cga gac tgt gga gat gct gcc cag caa ggc cgg gat ctg gcg cgt gga gtg cct gat cgg cga gca cct gca (SEQ ID NO:95)

23 r1

ggg gat cet cae gte tea get gge eat gee
cag ggg ggt etg gea ett gtt get gta eac
cag gaa eag ggt get eat gee gge gtg eag
gtg ete gee gat ea (SEQ ID NO:96)

23 bam

rl bbs 24 for (cagc)

cgc gaa ttc gga aga ccc cag cgg cca cat ccg cga ctt cca gat cac cgc cag cgg cca gta cgg cca gtg ggc tcc caa gct ggc ccg cct gca cta cag cgg (SEQ ID NO:97)

24 rl

ggg gat cet cae gte tea cat ggg gge cag

cag gte cae ett gat eea gga gaa ggg ete
ett ggt ega eea gge gtt gat get gee get
gta gtg eag geg gg (SEQ ID NO:98)

24 bam

5

r1 bbs 25 for (catg)

cgc gaa ttc gga aga ccc cat gat cat cca cgg cat caa gac cca ggg cgc ccg cca gaa gtt cag cag cct gta cat cag cca gtt cat cat cat gta ctc tct (SEQ ID NO:99) 25 rl

ggg gat cct cac gtc tca gtt gcc gaa gaa cac cat cag ggt gcc ggt gct gtt gcc gcg gta ggt ctg cca ctt ctt gcc gtc tag aga gta cat gat gat ga (SEQ ID NO:100)

25 bam

r1 bbs 26 for (caac)

cgc gaa ttc gga aga ccc caa cgt gga cag cag cgg cat caa gca caa cat ctt caa ccc ccc cat cat cgc ccg cta cat ccg cct gca ccc cac cca cta cag (SEQ ID NO:101) 26 rl

ggg gat cet cae gte tea gee eag ggg eat
get gea get gtt eag gte gea gee eat eag
ete eat geg eag ggt get geg gat get gta
gtg ggt ggg gtg ea (SEQ ID NO:102)

26 bam

rl bbs 27 for (gggc)

caa ggc cat cag cga cgc cca gat cac cgc ctc cag cta ctt cac caa cat gtt cgc cac ctg gag ccc cag caa (SEQ ID NO:103)

27 r1

ggg gat cet cae gte tea cea ete ett ggg gtt gtt cae etg ggg geg cea gge gtt get geg gee etg cag gtg cag geg gge ett get ggg get cea ggt gg (SEQ ID NO:104) 27 bam

5 r1 bbs 28 for (gtgg)

cgc gaa ttc gga aga ccc gtg gct gca ggt gga ctt cca gaa aac cat gaa ggt gac tgg cgt gac cac cca ggg cgt caa gag cct gct gac cag cat gta cgt (SEQ ID NO:105) 28 rl

ggg gat cet cae gte tea ett gee gtt ttg
gaa gaa eag ggt eea etg gtg gee gte etg
get get get gat eag gaa ete ett eae gta
eat get ggt eag ea (SEQ ID NO:106)

28 bam

rl bbs 29 for (caag)

gtt cca ggg caa cca gga cag ctt cac acc ggt cgt gaa cag cct gga ccc ccc cct gct gac ccg cta cct gcg (SEQ ID NO:107)

29 rl

ggg gat cet cae gte tea geg gee get tea

20 gta cag gte etg gge ete gea gee cag cae
cte cat geg cag gge gat etg gtg cae cea
get etg ggg gtg gat geg cag gta geg ggt
cag ca (SEQ ID NO:108)

29 bam

The codon usage for the native and synthetic genes described above are presented in Tables 3 and 4, respectively.

TABLE 3: Codon Frequency of the Synthetic Factor VIII B Domain Deleted Gene

5 .	AA	Codon	Numbe	r /100	0 Frac	ction
	Gly	GGG	7.00	4.82	0.09	
•	Gly	GGA	1.00	, 0.69	0.01	
	Gly	GGT	0.00	0.00	0.00	
10	Gly	GGC	74.00	50.93	0.90	٠,
	Glu	GAG	81.00	55.75	0.96	
	Glu	GAA	3.00	2.06	0.04	.1
	Asp	GAT ·	4.00	2.75	0.05	
15	Asp	GAC	78.00	53.68	0.95	•
	Val	GTG .	77.00	52.99	0.88	
	Val	GTA	2.00	1.38	0.02	
	Val	GTT	2.00	1.38	0.02	
20	Val	GTC ·	7.00	4.82	0.08	
	Ala	GCG	0.00	0.00	0.00	
	Ala	GCA	0.00	0.00	0.00	
	Ala	GCT	3.00	2.06	0.04	
25	Ala	GCC	67.00	46.11	0.96	
	Arg	AGG	2.00	1.38	0.03	
	Arg	AGA	0.00	0.00	0.00	
	Ser	AGT	0.00	0.00	0.00	
30	Ser	AGC	97.00	66.76	0.81	
	Lys	AAG	75.00	51.62	0.94	
	Lys	AAA	5.00	3.44	0.06	
	Asn	AAT	0.00	0.00	0.00	
35	Asn	AAC	63.00	43.36	1.00	

	Met	ATG	43.00	29.59	1.00
	Ile	AŢĀ	0.00	0.00	0.00
	Ile	ATT	2.00	1.38	0.03
5	Ile	ATC	72.00	49.55	0.97
	Thr	ACG	2.00	1.38	0.02
	Thr	AC'A	1.00	0.69	0.01
	Thr	ACT	10.00	6.88	0.12
10	Thr	ACC	70.00	48.18	0.84
•	Trp	TGG	28.00	19.27	1.00
	End	TGA	1.00	∗ 0.69	1.00
	Cys	TGT	1.00,	0.69	0.05
15	Cys	TGC	18.00	12.39	0.95
	End	TAG	0.00	0.00	0.00
	End	TAA	0.00	0.00	0.00
	Tyr	TAT	2.00	1.38	0.03
20	Tyr	TAC	66.00	45.42	0.97
	Leu	TTG	0.00	0.00	0.00
	Leu	TTA	0.00	0.00	0.00
	Phe	TTT	1.00	0.69	0.01
25	Phe	TTC	76.00	52.31	0.99
	Ser	TCG	1.00	0.69	0.01
	Ser	TCA	0.00	0.00	0.00
	Ser	TCT	-3.00	2.06	0.03
30	Ser	TCC	19.00	13.08	0.16
	Arg	CGG	1.00	0.69	0.01
	Arg	CGA	0.00	0.00	0.00
	Arg	CGT	1.00	0.69	0.01
35	Arg	CGC	69.00	47.49	0.95
	Gln	CAG	62.00	42.67	0.93
	Gln	CAA	5.00	3.44	0.07
	His	CAT	1.00	0.69	0.02
40	His	CAC	50.00	34.41	0.98

·	Leu Leu	CTG CTA	118.00	81:2 <u>1</u> 2.06	0.94 0.02
5	Leu Leu	CTC	1.00 3.00	0.69 2.06	0.01
10	Pro Pro Pro Pro	CCG CCA CCT CCC	4.00 0.00 3.00 68.00	2.75 0.00 2.06 46.80	0.05 0.00 0.04 0.91

TABLE 4: Codon Frequency Table of the Native Factor VIII B Domain Deleted Gene

15 '	AA	Codon	Numb	per /100	00 Fraction	
	Gly	GGG	12.00	8.26	0.15	
	Gly	GGA	34.00	23.40	0.41	
	Gly	GGT	16.00	11.01	0.20	
20	Gly	GGC	20.00	13.76	0.24	
	Glu	GAG	33.00	22.71	0.39	
	Glu	GAA	51.00	35.10	0.61	
	Asp	GAT	55.00	37.85	0.67	
25	Asp	GAC	27.00	18.58	0.33	
	Val	GTG	29.00	19.96	0.33	
	Val	_GTA	_19.00_	-13.08	-0.22	
	Val	GTT	17.00	11.70	0.19	
30	Val	GTC	23.00	15.83	0.26	
	Ala	GCG	2.00	1.38	0.03	
	Ala	GCA	18.00	12.39	0.25	
	Ala	GCT	31.00	21.34	0.44	
35	Ala	GCC	20.00	13.76	0.28	

	Arg	AGG	18.00	12.39	0.25
	Arg		22.00	15.14	
	Ser				0.18
	Ser	AGC	24.00		
5					. 9,20
	Lys	AAG	32.00	22.02	0.40
	Lys	AAA	48.00	33.04	0.60
	Asn	AAT	38.00	26.15	0.60
•	Asn	AAC	25.00	17.21	0.40
10	•		•		
	Met	ATG	43.00	29.59	1.00
	Ile	ATA	13.00	8.95	0.18
	Ile	ATT	36.00	24.78	0.49
	Ile	ATC	25.00	17.21	0.34
15		1	•		
	Thr	ACG	1.00	0.69	0.01
	Thr	ACA	23.00	15.83	0.28
	Thr	ACT	36.00	24.78	0.43
	Thr	ACC	23.00	15.83	0.28
20			I		
	Trp	TGG	28.00	19.27	1.00
	End	TGA	1.00	0.69	1.00
	Cys	TGT	7.00 .	4.82	0.37
	Cys	TGC	12.00	8.26	0.63
25		,			
	End	TAG	0.00	0.00	0.00
	End	TAA	0.00	0.00	0.00
	Tyr	TAT	41.00	28.22	0.60
	Tyr	TAC	27.00	18.58	0.40
30					
	Leu	TTG	20.00	13.76	0.16
	Leu	TTA	10.00	6.88	0.08
	Phe	TTT	45.00	30.97	0.58
	Phe	TTC	32.00	22.02	0.42
35					
•	Ser	TCG	2.00	1.38	0.02
	Ser	TCA	27.00	18.58	0.22
	Ser	TCT	27.00	18.58	0.22
	Ser	TCC	18.00	12.39	0.15
40					

						1
1	Arg	CGG	6.00	4.13	0.08	· · · · · · · · · · · · · · · · · · ·
	Arg	CGA	10.00	6.88	0.14	•
	Arg	CGT	7.00	4.82,	0.10	
	Arg	CGC	10.00	6.88	0.14	() () () () () () () () () ()
5						
	Gln	CAG	42.00	28.91	0.63	
·	Gln	CAA	25.00	17.21	0:37	
	His	CAT	28.00	19.27	0.55	1
	His	CAC	23.00	15.83	0.45	
10		,				•
	Leu	CTG	36.00	24.78	0.29	•
. '	Leu	CTA -	15.00	10.32	0.12	
	Leu	CTT	24.00	16.52	0.19	
	Leu	CTC	20.00	13.76	0.16	·
15				:		
1	Pro	CCG	1.00	0.69	0.01	ı
	Pro	CCA	32.00	22.02	0.43	
ı	Pro	CCT	26.00	17.89	0.35	
	Pro	· CCC	15.00	10.32	0.20	
20						

<u>Use</u>

25

30

The synthetic genes of the invention are useful for expressing the a protein normally expressed in mammalian cells in cell culture (e.g. for commercial production of human proteins such as hGH, TPA, Factor VIII, and Factor IX). The synthetic genes of the invention are also useful for gene therapy. For example, a synthetic gene encoding a selected protein can be introduced in to a cell which can express the protein to create a cell which can be administered to a patient in need of the protein. Such cell-based gene therapy techniques are well known to those skilled in the art, see, e.g., Anderson, et al., U.S. Patent No. 5,399,349; Mulligan and Wilson, U.S. Patent No. 5,460,959.

What is claimed is:

1. A synthetic gene encoding a protein normally expressed in an eukaryotic cell wherein at least one non-preferred or less preferred codon in a natural gene encoding said protein has been replaced by a preferred codon encoding the same amino acid, said synthetic gene being capable of expressing said protein at a level which is at least 110% of that expressed by said natural gene in an *in vitro* mammalian cell culture system under identical conditions.

- 2. The synthetic gene of claim 1 wherein said synthetic gene is capable of expressing said protein at a level which is at least 150% of that expressed by said natural gene in an *in vitro* cell culture system under identical conditions.
- 3. The synthetic gene of claim 1 wherein said synthetic gene is capable of expressing said protein at a level which is at least 200% of that expressed by said natural gene in an *in vitro* cell culture system under identical conditions.
- 4. The synthetic gene of claim 1 wherein said synthetic gene is capable of expressing said protein at a level which is at least 500% of that expressed by said natural gene in an *in vitro* cell culture system under identical conditions.
- 5. The synthetic gene of claim 1 wherein said synthetic gene comprises fewer than 5 occurrences of the sequence CG.
 - 6. The synthetic gene of claim 1 wherein at least 10% of the codons in said natural gene are non-preferred codons.

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7. The synthetic gene of claim 1 wherein at least 50% of the codons in said natural gene are non-preferred codons.

- 8. The synthetic gene of claim 1 wherein at least 50% of the non-preferred codons and less preferred codons present in said natural gene have been replaced by preferred codons.
- 9. The synthetic gene of claim 1 wherein at least 90% of the non-preferred codons and less preferred codons present in said natural gene have been replaced by preferred codons.
- 10. The synthetic gene of claim 1 wherein said protein is normally expressed by a mammalian cell.
 - 11. The synthetic gene of claim 1 wherein said protein is a retroviral protein.
 - 12. The synthetic gene of claim 1 wherein said protein is a lentiviral protein.
- 13. The synthetic gene of claim 11 wherein said protein is an HIV protein.
 - 14. The synthetic gene of claim 13 wherein said protein is selected from the group consisting of gag, pol, and env.
 - 15. The synthetic gene of claim 13 wherein said protein is gp120.

16. The synthetic gene of claim 13 wherein said protein is gp160.

- 17. The synthetic gene of claim 1 wherein said protein is a human protein.
- 18. The synthetic gene of claim 1 wherein said human protein is Factor VIII.
 - 19. The synthetic gene of claim 1 wherein 20% of the codons are preferred codons.
 - 20. The synthetic gene of claim 18 wherein said gene has the coding sequence present in SEQ ID NO:42.
- The synthetic gene of claim 1 wherein said protein is green fluorescent protein.
 - 22. The synthetic gene of claim 20 wherein said synthetic gene is capable of expressing said green fluorescent protein at a level which is at least 200% of that expressed by said natural gene in an *in vitro* mammalian cell culture system under identical conditions.
 - 23. The synthetic gene of claim 20 wherein said synthetic gene is capable of expressing said green fluorescent protein at a level which is at least 1000% of that expressed by said n atural gene in an *in vitro* mammalian cell culture system under identical conditions.

24. The synthetic gene of claim 21 having the sequence depicted in Figure 11 (SEQ ID NO:40).

- 25. An expression vector comprising the synthetic gene of claim 1.
- 5 26. The expression vector of claim 21, said expression vector being a mammalian expression vector.
 - 27. A mammalian cell harboring with the synthetic gene of claim 1.
- 28. A method for preparing a synthetic gene encoding a protein normally expressed by mammalian cells, comprising identifying non-preferred and less-preferred codons in the natural gene encoding said protein and replacing one or more of said non-preferred and less-preferred codons with a preferred codon encoding the same amino acid as the replaced codon.

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· Syngpl20mn

1 CTCGAGATCC ATTGTGCTCT AAAGGAGATA CCCGGCCAGA CACCCTCACC TGCGGTGCCC AGCTGCCCAG GCTGAGGCAA GAGAAGGCCA GAAACCATGC CCATGGGGTC TETGCAACCG CTGGCCACCT TGTACCTGCT GGGGATGCTG 101 151 GTCGCTTCCG TGCTAGCCAC CGAGAAGCTG TGGGTGACCG TGTACTACGG 201 CGTGCCCGTG TEGAAGGAGG CCACCACCAC CCTGTTCTGC GCCAGCGACG 251 CCAAGGCGTA CBACACCGAG GTGCACAAGG TGTGGGCCAC CCAGGCGTGC GTGCCCACCG ACCCCAACCC CCAGGAGGTG GAGCTCGTGA ACGTGACCGA GAACTTCAAC NEGTGGAAGA ACAACATGGT GGAGCAGATG CATGAGGACA TCATCAGCCT GTGGGACCAG AGCCTGAAGC CCTGCGTGAA GCTGACCCCC CTGTGCGTGA (ECTGAACTG CACCGACCTG AGGAACACCA CCAACACCAA CAACAGCACC GCCAACAACA ACAGCAACAG CGAGGGCACC ATCAAGGGCG GCGAGATGAA CAACTGCAGC TTCAACATCA CCACCAGCAT CCGCGACAAG ATGCAGAAGG ASTACGCCCT GCTGTACAAG CTGGATATCG TGAGCATCGA CAACGACAGC ACCAGCTACC GCCTGATCTC CTGCAACACC AGCGTGATCA CCCAGGCCTG GCCCAAGATC AGCTTCGAGC CCATCCCCAT CCACTACTGC GCCCCCCCC CCTTCGCCAT CCTGAAGTGC AACGACAAGA AGTTCAGCGG CAAGGGCAGC TGCAAGAACG TGAGCACCGT GCAGTGCACC CACGGCATCC GGCCGGTGGT GAGCACCCAG CTCCTGCTGA ACGGCAGCCT GGCCGAGGAG GAGGTGGTGA TCCGCAGCGA GAACTTCACC GACAACGCCA AGACCATCAT CGTGCACCTG AATGAGAGCG TGCAGATCAA CTGCACGCGT CCCAACTACA ACAAGCGCAA GCGCATCCAC ATCGGCCCCG GGCGCGCCTT CTACACCACC AAGAACATCA TCGGCACCAT CCGCCAGGCC CACTGCAACA TCTCTAGAGC CAAGTGGAAC GACACCCTGC GCCAGATCGT GAGCAAGCTG AAGGAGCAGT TCAAGAACAA GACCATCGTG TTCAACCAGA GCAGCGGCGG CGACCCCGAG ATCGTGATGC ACAGCTTCAA CTGCGGCGGC GAATTCTTCT ACTGCAACAC 1201 CAGCCCCTG TTCAACAGCA CCTGGAACGG CAACAACACC TGGAACAACA CCACCGGCAG CAACAACAAT ATTACCCTCC AGTGCAAGAT CAAGCAGATC ATCAACATGT GGCAGGAGGT GGGCAAGGCC ATGTACGCCC CCCCCATCGA GGGCCAGATC CGGTGCAGCA GCAACATCAC CGGTCTGCTG CTGACCCGCG 1451 ACOGCGGAA GGACACGAC ACCAACGACA CAGAAATCTT CCGCCCCCGC

> FIGI (SHEET 1 OF 4)

1501 GGGGGGACA TGCGCGACAA CTGGAGATCT GAGCTGTACA AGTACAAGGT
1551 GGTGACGATC GAGCCCCTGG GCGTGGCCCC CACCAAGGCC AAGCGCCGCG
1601 TGGTGCAGCG CGAGAAGCGC TAAAGCGGCC GC (SEQ ID NO:34)

FIG | (SHEET 2 OF 4)

syngploomn.

1 AUCUAGAAGO TOTGGGTGAC COTGTACTAC GGCGTGCCCG TGTGGAAGGA 51 GGCCACCACC MICCIGITCT GCGCCAGCGA CGCGAAGGCG TACGACACCG AUGTGCACAA CETGTGGGCC ACCCAGGCGT GCGTGCCCAC CGACCCCAAC CCCCAGGAGG TEGACCTCGT GAACGTGACC GAGAACTTCA ACATGTGGAA GAACAACATG CTGGAGCAGA TGCATGAGGA CATCATCAGC CTGTGGGACC 201 AGAGCCTGAA GOCCTGCCTG AAGCTGACCC CCCTGTGCGT GACCCTCAAC TGEACCGACC TGAGGAACAC CACCÁACACC AACAACAGCA CCGCCAACAA CHACAGCAAC AGCGAGGGCA CCATCAAGGG CGGCGAGATG AAGAACTGCA GOTTCAACAT CACCACCAGC ATCCGCGACA AGATCCAGAA GGAGTACGCC CTGCTGTACA AGCTGGATAT CGTGAGCATC CACAACGACA GCACCAGCTA 501 CCGCCTGATC TCCTGCAACA CCAGCGTGAT CACCCAGGCC TGCCCCAAGA TCAGCTTCGA GCCCATCCCC ATCCACTACT GCGCCCCCGC CGGCTTCGCC ATCCTGAAGT QCAACGACAA GAAGTTCAGC GGCAAGGGCA GCTGCAAGAA CGTGACCACC CTGCAGTGCA CCCACGGCAT CCGGCCGGTG GTGAGCACCC ACCTCCTGCT GAACGGCAGC CTGGCCGAGG AGGAGGTGGT GATGGGCAGC GAGAACTIVA CCGACAACGC CAAGACCATC ATCGTGCACC TGAATGAGAG CGTGCAGATC AACTGCACGC GTCCCAACTA CAACAAGCGC AAGCGCATCC 851 ACATOGOCCO COGGOGGGCC TTCTACACCA CCAAGAACAT CATCGGCACC 901 ATCCGCCAGG CCCACTGCAA CATCTCTAGA GCCAAGTGGA ACGACACCCT 951 GCGCCAGATC GTGAGCAAGG TGAAGGAGCA GTTCANGAAC AAGACCATCG TGTTCAACCA GAGCAGCGGC GGCGACCCCG AGATCGTGAT GCACAGCTTC AACTGCGGGG GCGAATTCTT CTACTGCAAC ACCAGCCCCC TGTICAACAG CACCTGGAAC GGCAACAACA CCTGGAACAA CACCACUGGC AGCAACAACA ATATTACCCT CCAGTGCAAG ATCAAGCAGA TCATCAACAT GTGGCAGGAG 1201 GTGGGCAAGG CCATGTACGC CCCCCCATC GAGGGCCAGA TCCGGTGCAG 1251 CAGCAACATO ACCOUTUTGO TOCTGACCOS CGACGGCGGC AAGGACACCG 1301 ACACCANCUA CACCGAAATC TTCCGCCCCG GCGGGGGGGGA CATGCGCGAC AAUTGGAGAT CTGAGCTGTA CAAGTACAAG GTGGTGACGA TCGAGCCCCT COCCOTOCCC CCCACCAAGG CCAAGGGCCG CGTGGTGCAG CGCGAGAAGC

> FIG. 1 (SHEET 3 OF 4)

1451	GGGCCGCCAT CUGCGCCCTG TTCCTGGGCT TCCTGGGGGC GGCGGCAGC
1501	ACCATGGGGG CCGCCAGCGT GACCCTGACC GTGCAGGCCC GCCTGCTCCT
1551	GAGCGGCATC GTGCAGCAGC AGAACAACCT CCTCCGCGCC ATCGAGGCCC
1601	AGCAGCATAT GITCCAGCTC ACCGTGTGGG GCATCAAGCA GCTCCAGGCC
1651	CGCGTGCTGG CCGTGGAGCG CTACCTGAAG GACCAGCAGC TCCTGGGCTT
.1701	CTGGGGCTGC TCCGGCAAGC TGATCTGCAC CAGGACGGTA CCCTGGAACG
1751	CCTCCTGGAG CAACAAGAGC CTGGACGACA TCTGGAACAA CATGACCTGG
1501	ATGCAGTGGG AGCGCGAGAT CGATAACTAG ACCAGCCTGA TCTACAGCCT
1851	GCTGGAGAAG AACCAGACCC AGCAGGAGAA GAACGAGCAG GAGCTGCTGG
1901	AGGTGGACAA CIGGGCGAGC CTGTGGAACT GGTTCGACAT CACCAACTGG
1951	CTGTGGTACA TCAAAATCTT CATCATGATT GTGGGCGGCC TGGTGGGCCT
2001	CCGCATCGTG TTCGCCGTGC TGAGCATCGT GAACCGCGTG CGCCAGGGCT
2051	ACAGCCCCT GAGCCTCCAG ACCCGGCCCC CCGTGCCGCG CGGGCCCGAC
2101	CGCCCCGAGG CCATCGAGGA GGAGGGCGGC GAGCGCGACC GCGACACCAG
2151	CGGCAGGCTC GTGCACGGCT TCCTGGCGAT CATCTGGGTC GACCTCCGCA
2201	SCCTGTTCCT GTTCAGCTAC CACCACCGCG ACCTGCTGCT GATCGCCGCC
2251	CGCATCGTGG AACTCCTAGG CCGCCGCGC TGGGAGGTGC TGAAGTACTG
2301	GTGGAACCTC CTCCAGTATT GGAGCCAGGA GCTGAAGTCC AGCGCCGTGA
2351	GCCTGCTGAA CGCCACCGCC ATCGCCGTGG CCGAGGGCAC CGACCGCGTG
2401	ATCGAGGTGC TCCAGAGGGC CGGGAGGGCG ATCCTGCACA TCCCCACCCG
2451	CATCCGCCAG (GGCTCGAGA GGGCGCTGCT G (SEQ ID NO:35)

FIG. 1 (SHEET 4 OF 4)

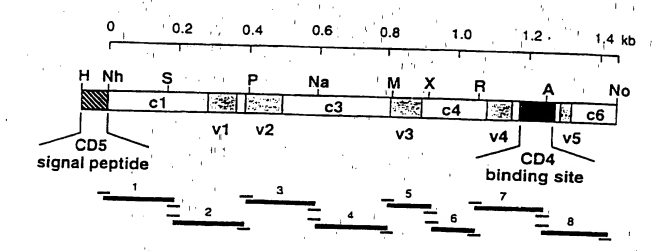


FIGURE 2

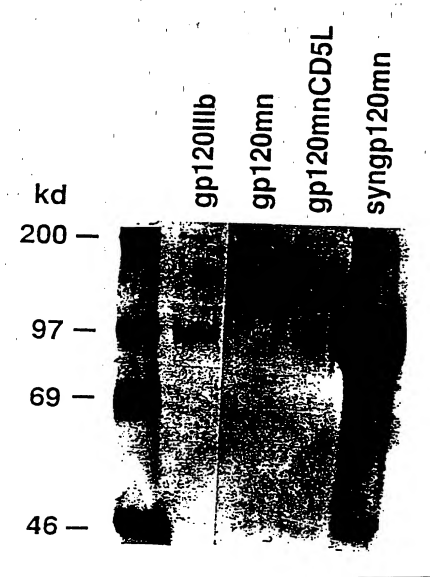


FIGURE 3

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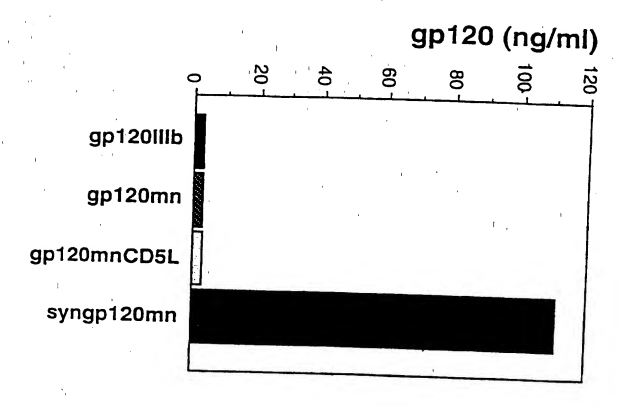


FIGURE 4

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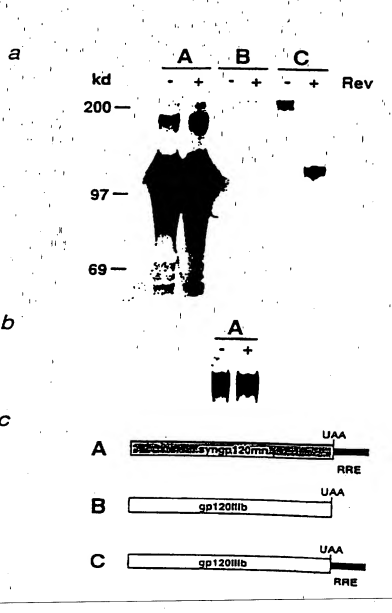


FIGURE 5

Q Caa cag act ပ agt tct K · K K T I ຸຜ aac aat gag aat ata ·田 · O aac Z att C Lgt × aga gta gtc gat gat gaa aaa aaa œ caa aaa > caa ttt L D tta gat 1 gac zaa_ agt tcc aga េ gag aag T D aca^{gat} Cgc aat tta gta gac gtt > ctg F T T ttt aca aca R Cgť agt tcc atg atg cga S agt tcc . .__ acc ctg Σ T acg / R aga cga caa P T S Cca aca_agt acc cag R aga cgc acc agc đ tta caa gca gcc ctg J K L ttg ctt r tta ctg-L tta ttg Y tat tac Ţ aca aac ttc S agt agc caa Ö Q N caa aat aac F S ttt tca t ttc ago c gta gtc aca N aat act ၁၁၁ ata tta ata ctc H Δ, H cag L S tta agt tca aat A gca gcc cac н Cat gga aat ttt agc တ္ CL, gta aat gtg aa E gaa gag T L aca tta ctt E gaa gag z cta Caa cag gga s agt tcc ggt G gtg tta ctg H Cat . E act. cca cat ၁၁၁ 99a 99c L tta ctc ۵, C tgt tgt ပ C L tgt tta tgc ctg tta cca ata caa gta tta gtc ctt gta cag gtt s agt tcg ж ааа aag s agt tcc ø > J ID NO:36) env-batg aat cca gta ata agt ata aca ID NO:37)-wt--> atg aac cca gtc atc agc atc act atc gta gtc gga 666 gta Н L tta ctt · છ > R V I S L T A aga gta ata agt tta aca gca agg gtg atc agc ctg aca gcc ၁၁၁ L tta ctg aaa ۵, aag aga cga L tta ctg L tta ctg × **~** L ttg ata aca acc ctc ctt aaa aag S L ta Н L N T N aat aca cct t aac acc aac t G gga ggc tto gag gaa gat gac Œ L tta ctg ш Ω aga cgc aga. aga s agt tca tta c tgt tgt œ α; tta gat gac ctg atg atg ata tta Q ctg Σ **-**ᄀ aat .v gta gtg aat agt agt Y tat tac gta gtg tgg tgg z S > gaa R Cat ttt D gat gac aat S agt tcc Z env Wt env env wt env wt env WC env wt env wt Σ

ICURE 6

tga tga

tta ctg

env wc

(SEQ (SEQ 10/18

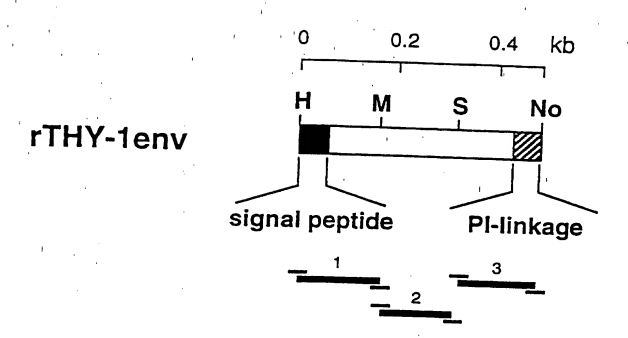


FIGURE 7

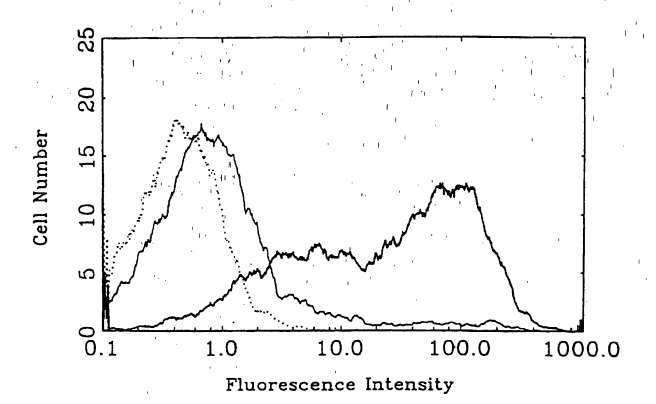
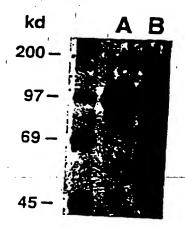


FIGURE 8

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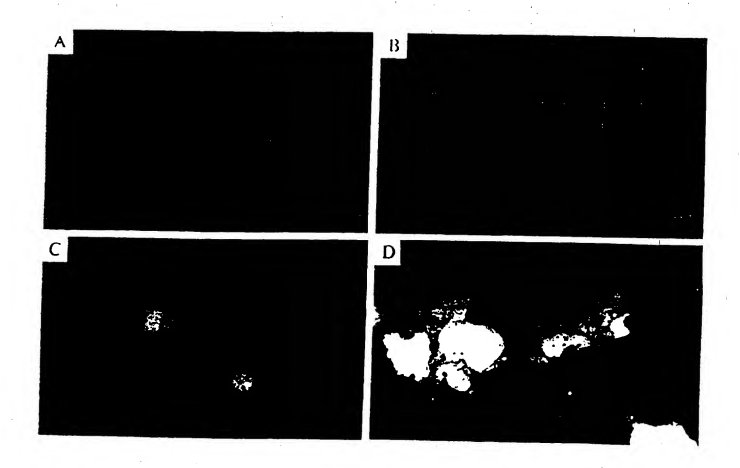
b

A syngp120mn

B syngp120min rTHY-1env

FIGURE 9

FIG. 10



1	GAATTCACGC GTAAGCTTGC CGCCACCATG GTGAGCAAGG GCGAGGAGC
51	GTTCACCGGG GTGGTGCCCA TCCTGGTCGA GCTGGACGGC GACGTGAAC
101	GCCACAAGTT CAGCGTGTCC GGCGAGGGCG AGGGCGATGC CACCTACGG
151	AAGCTGACCC TGAAGTTCAT CTGCACCACC GGCAAGCTGC CCGTGCCCTG
201	GCCCACCTC GTGACCACCT TCAGCTACGG CGTGCAGTGC TTCAGCCGCT
251	ACCCCGACCA CATGAAGCAG CACGACTTCT TCAAGTCCGC CATGCCCGAA
301	GGCTACGTCC AGGAGCGCAC CATCTTCTTC AAGGACGACG GCAACTACAA
351	GACCCGCGCC GAGGTGAAGT TCGAGGGCGA CACCCTGGTG AACCGCATCG
401	AGCTGAAGGG CATCGACTTC AAGGAGGACG GCAACATCCT GGGGCACAAG
151	CTGGAGTACA ACTACAACAG CCACAACGTC TATATCATGG CCGACAAGCA
501	GAAGAACGGC ATCAAGGTGA ACTTCAAGAT CCGCCACAAC ATCGAGGACG
551	GCAGCGTGCA GCTCGCCGAC CACTACCAGC AGAACACCCC CATCGGCGAC
01	GGCCCCGTGC TGCTGCCCGA CAACCACTAC CTGAGCACCC AGTCCGCCCT
51	GAGCAAAGAC CCCAACGAGA AGCGCGATCA CATGGTCCTG CTGGAGTTCG
01	TGACCGCCGC CGGGATCACT CACGGCATGG ACGAGCTGTA CAAGTAAAGC
51	GGCCGCGGAT CC (SEC ID NO. 40)

Native Factor VIII B domain deleted gene segment inserted in the expr ssion vector

	,				
1	AAGCTTAAAC	CATGCCCATG	GGGTCTCTGC	AACCGCTGGC	CACCTTGTAC
51	CTGCTGGGGA	TGCTGGTCGC	TTCCGTGCTA	GCCGCCACCA	GAAGATACTA
101	CCTGGGTGCA	GTGGAACTGT	CATGGGACTA	TATGCAAAGT	GATCTCGGTG
151	AGCTGCCTGT	GGACGCAAGA	TTTCCTCCTA	GAGTGCCAAA	ATCTTTTCCA
201	TTCAACACCT	CAGTCGTGTA	CAAAAAGACT	CTGTTTGTAG	AATTCACGGA
251		AACATCGCTA			GGTCTGCTAG
301	GTCCTACCAT	CCAGGCTGAG	GTTTATGATA	CAGTGGTCAT	TACACTTAAG
351	AACATGGCTT	CCCATCCTGT	CAGTCTTCAT	GCTGTTGGTG	TATCCTACTG
401	GAAAGCTTCT	GAGGGAGCTG	AATATGATGA	TCAGACCAGT	CAAAGGGAGA
451	A'AGAAGATGA	TAAAGTCTTC	CCTGGTGGAA	GCCATAÇATA	TGTCTGGCAG
501	GTCCTGAAAG	AGAATGGTCC	AATGGCCTCT	GACCCACTGT	GCCTTACCTA
551	CTCATATCTT	TCTCATGTGG	ACCTGGTAAA	AGACTTGAAT	TCAGGCCTCA
601	TTGGAGCCCT	ACTAGTATGT	AGAGAAGGGA	GTCTGGCCAA	GGAAAAGACA
651	CAGACCTTGC		ACTACTTTTT		ATGAAGGGAA
701		TCAGAAACAA	-	GATGCAGGAT	AGGGATGCTG
751	CATCTGCTCG			CAGTCAATGG	TTATGTAAAC
801	AGGTCTCTGC	CAGGTCTGAT	TGGATGCCAC	AGGAAATCAG	TCTATTGGCA
851	TGTGATTGGA	ATGGGCACCA		GCACTCAATA	TTCCTCGAAG
901	GTCACACATT			AGGCGTCCTT	GGAAATCTCG
951			TCAAACACTC	TTGATGGACC	TTGGACAGTT
1001	TCTACTGTTT		CTTCCCACCA		ATGGAAGCTT
1051	ATGTCAAAGT	AGACAGCTGT	CCAGAGGAAC	CCCAACTACG	AATGAAAAAT
1101	AATGAAGAAG		TGATGATGAT	CTTACTGATT	CTGAAATGGA
1151	TGTGGTCAGG	TTTGATGATG	ACAACTCTCC		CAAATTCGCT
1201		GAAGCATCCT	AAAACTTGGG	TACATTACAT	TGCTGCTGAA
1251	GAGGAGGACT	GGGACTATGC			ATGACAGAAG
1301	TTATAAAAGT	CAATATTTGA		TCAGCGGATT	GGTAGGAAGT
1351	ACAAAAAGT	1		ATGAAACCTT	TAAGACTCGT
1401	GAAGCTATTC	AGCATGAATC		GGACCTTTAC	TTTATGGGGA
1451		ACACTGTTGA	TTATATTTAA		AGCAGACCAT
1501	ATAACATCTA	CCCTCACGGA		TCCGTCCTTT	GTATTCAAGG
1551		•	ACATTTGAAG	GATTTTCCAA	
1601		AAATATAAAT	GGACAGTGAC	TGTAGAAGAT	GGGCCAACTA
1651	AATCAGATCC.		ACCCGCTATT	ACTCTAGTTT	CGTTAATATG
1701			ACTCATTGGC		TCTGCTACAA
1751		GATCAAAGAG	GAAACCAGAT		AAGAGGAATG
1801	TCATCCTGTT	TTCTGTATTT	GATGAGAACC	GAAGCTGGTA	CCTCACAGAG
1851		GCTTTCTCCC		GGAGTGCAGC	TTGAGGATCC TATGTTTTTG
1901 1951	AGAGTTCCAA ATAGTTTGCA	GCCTCCAACA GTTGTCAGTT	TGTTTGCATG	CATCAATGGC AGGTGGCATA	CTGGTACATT
2001	CTAAGCATTG		TGACTTCCTT	TCTGTCTTCT	TCTCTGGATA
2051	TACCTTCAAA	CACAAAATGG	TCTATGAAGA	CACACTCACC	CTATTCCCAT
2101	TCTCAGGAGA	AACTGTCTTC	ATGTCGATGG	AAAACCCAGG	TCTATGGATT
2151		ACAACTCAGA			
2201	GAAGGTTTCT		AGAACACTGG	TGATTATTAC	
2251	ATGAAGATAT	TTCAGCATAC		AAAACAATGC	
2301	AGAAGCTTCT			AGCACTAGGC	
2351	TAATGCCACC	CCACCAGTCT	TGAAACGCCA	TCAACGGGAA	
2401	CTACTCTTCA		GAGGAAATTG	ACTATGATGA	
2451	GTTGAAATGA		TTTTGACATT	TATGATGAGG	
2501	GAGCCCCCGC	AGCTTTCAAA	AGAAAACACG	ACACTATTTT	ATTGCTGCAG
2551	TGGAGAGGCT	CTGGGATTAT	GGGATGAGTA	GCTCCCCACA	TGTTCTAAGA
2601		AGAGTGGCAG	TGTCCCTCAG	TTCAAGAAAG	TTGTTTTCCA
2651		GATGGCTCCT		CTTATACCGT	
2701				TAAGAGCAGA	
	·				

Fig. 12

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				•		
:	2751	AATATCATGG	TAACTTTCAG	AAATCAGGCC	TCTCGTCCCT	ATTCCTTCTA
	2801	TTCTAGCCTT	ATTTCTTATG	AGGAAGATCA	GAGGCAAGGA	GCAGAACCTA
٠.	2851	GAAAAAACTT	TGTCAAGCCT	AATGAAACCA	AAACTTACTT	TTGGAAAGTG
:	2901	CAACATCATA	TGGCACCCAC	TAAAGATGAG	TTTGACTGCA	AAGCCTGGGC
	2951	TTATTTCTCT	GATGTTGACC	TGGAAAAAGA	TGTGCACTÇA	GCCTGATTG
	3001	GACCCCTTCT	GGTCTGCCAC	ACTAACACAC	TGAACCCTGC	TCATGGGAGA
	3051	CAAGTGACAG	TACAGGAATT	TCCTCTGTTT	TTCACCATCT	TTGATGAGAC
	3101	CAAAAGCTGG	TACTTCACTG	AAAATATGGA	AAGAAACTGC	AGGGCTCCCT
	3151	GCAATATCCA	GATGGAAGAT	CCCACTTTTA	AAGAGAATTA	TCGCTTCCAT
	3201	GCAATCAATG	GCTACATAAT	GGATACACTA	CCTGGCTTAG	TAATGGCTCA
	3251	GGATCAAAGG	ATTCGATGGT	ATCTGCTCAG	CATGGGCAGC	AATGAAAACA
	3301	TCCATTCTAT	TCATTTCAGT	GGACATGTGT	TCACTGTACG	AAAAAAAGAG
	3351	GAGTATAAAA	TGGCACTGTA	CAATCTCTAT	CCAGGTGTTT	TTGAGACAGT
	3401	GGAAATGTTA	CCATCCAAAG	CTGGAATTTG	GCGGGTGGAA	TGCCTTATTG
	3451	GCGAGCATCT	ACATGCTGGG	ATGAGCACAC	TTTTTCTGGT	GTACAGCAAT
	3501	AAGTGTCAGA	CTCCCCTGGG	AATGGCTTCT	GGACACATTA	GAGATTTTCA
	3551	GATTACAGCT	TCAGGACAAT	ATGGACAGTG	GGCCCCAAAG	CTGGCCAGAC
	3601	TTCATTATTC	CGGATCAATC	AATGCCTGGA	GCACCAAGGA	GCCCTTTTCT
i	3651	TGGATCAAGG	TGGATCTGTT	GGCACCAATG	ATTATTCACG	GCATCAAGAC
	3701	CCAGGGTGCC	CGTCAGAAGT	TCTCCAGCCT	CTACATCTCT	CAGTTTATCA
	3751	TCATGTATAG	TCTTGATGGG	AAGAAGTGGC	AGACTTATCG	AGGAAATTCC
	3801	ACTGGAACCT	TAATGGTCTT	CTTTGGCAAT	GTGGATTCAT	CTGGGATAAA
	3851	ACACAATATT	TTTAACCCTC	CAATTATTGC	TCGATACATC	CGTTTGCACC
	3901	CAACTCATTA	TAGCATTCGC	AGCACTCTTC	GCATGGAGTT	GATGGGCTGT
	3951	GATTTAAATA	GTTGCAGCAT	GCCATTGGGA		AAGCAATATC
	4001	AGATGCACAG	ATTACTGCTT	CATCCTACTT	TACCAATATG	TTTGCCACCT
	4051	GGTCTCCTTC	AAAAGCTCGA	CTTCACCTCC	AAGGGAGGAG	TAATGCCTGG
	4101	AGACCTCAGG	TGAATAATCC	AAAAGAGTGG	CTGCAAGTGG	ACTTCCAGAA
	4151	GACAATGAAA	GTCACAGGAG	TAACTACTCA	GGGAGTAAAA	TCTCTGCTTA
	4201	CCAGCATGTA	TGTGAAGGAG	TTCCTCATCT	CCAGCAGTCA	AGATGGCCAT
	4251	CAGTGGACTC	TCTTTTTCA	GAATGGCAAA	GTAAAGGTTT	TTCAGGGAAA
	4301	TCAAGACTCC	TTCACACCTG	TGGTGAACTC	TCTAGACCCA	CCGTTACTGA
	4351	CTCGCTACCT	TCGAATTCAC	CCCCAGAGTT	GGGTGCACCA	GATTGCCCTG
	4401	AGGATGGAGG	TTCTGGGCTG	CGAGGCACAG	GACCTCTACT	GAGGGTGGCC
	4451	ACTGCAGCAC	CTGCCACTGC	CGTCACCTCT	CCCTCCTCAG	CTCCAGGGCA
	4501	GTGTCCCTCC	CTGGCTTGCC	TTCTACCTTT	GTGCTAAATC	CTAGCAGACA
	4551	CTGCCTTGAA	GCCTCCTGAA	TTAACTATCA	TCAGTCCTGC	ATTTCTTTGG
	4601	TGGGGGGCCA		TCCAATTTAA	CTTAACTCTT	ACCGTCGACC
•	4651	TGCAGGCCCA	ACGCGGCCGC			

Fig. 12

(2 of 2)

Synthetic Factor VIII B domain delet d gene segment inserted in the expression vector

```
1 AAGCTTAAAC CATGCCCATG GGGTCTCTGC AACCGCTGGC CACCTTGTAC
           CTGCTGGGGA TGCTGGTCGC TTCCGTGCTA GCCGCCACCC GCCGCTACTA
     51
           CCTGGCCCC GTGGAGCTGT CCTGGGACTA CATGCAGAGC GACCTGGGCG
AGCTCCCCGT GGACGCCCGC TTCCCCCCCCC GCGTGCCCAA GAGCTTCCCC
   101
    151
          TTCAACACCA GCGTGGTGTA CAAGAAAACC CTGTTCGTGG AGTTCACCGA
   201
           CCACCTGTTC AACATTGCCA AGCCGCGCCC CCCCTGGATG GGCCTGCTGG
   251
   301 GCCCACCAT CCAGGCCGAG GTGTACGACA CCGTGGTGAT CACCCTGAAG
351 AACATGGCCA GCCACCCCGT CAGCCTGCAC GCCGTGGGCG TGAGCTACTG
   401 GAAGGCCAGC GAGGGCGCCG AGTACGACGA CCAGACGTCC CAGCGCGAGA
   AGGAGACGA CAAGGTGTTC CCGGGGGGGA GCCACACCTA CGTGTGGCAG
501 GTGCTTAAGG AGAACGGCCC TATGGCCAGC GACCCCCTGT GCCTGACCTA
551 CAGCTACCTG AGCCACGTGG ACCTGGTGAA GGATCTGAAC AGCGGGCTGA
601 TCGGCGCCCT GCTGGTGTT CGCGAGGGCA GCCTGGCCAA GGAGAAAACC
         CAGACCCTGC ACAAGTTCAT CCTGCTGTTC GCCGTGTTCG ACGAGGGGAA
   651
   701
          GAGCTGGCAC AGCGAGACTA AGAACAGCCT GATGCAGGAC CGCGACGCCG
   751
         CCAGCGCCCG CGCCTGGCCC AAGATGCACA CCGTTAACGG CTACGTGAAC
         CGCAGCCTGC CCGGCCTGAT CGGCTGCCAC CGCAAGAGCG TGTACTGGCA
CGTCATCGGC ATGGGCACCA CCCCTGAGGT GCACAGCATC TTCCTGGAGG
   801
   851
   901
         GCCACACCTT CCTGGTGCGC AACCACCGCC AGGCCAGCCT GGAGATCAGC
 951 CCCATCACCT TCCTGACTGC CCAGACCCTG CTGATGGACC TAGGCCAGTT
1001 CCTGCTGTTC TGCCACATCA GCAGCCACCA GCACGACGGC ATGGAGGCTT
1051 ACGTGAAGGT GGACAGCTGC CCCGAGGAGC CCCAGCTGCG CATGAAGAAC
1101 AACGAGGAGG CCGAGGACTA CGACGACGAC CTGACCGACA GCGAGATGGA
 1151 TGTCGTACGC TTCGACGACG ACAACAGCCC CAGCTTCATC CAGATCCGCA
1201 GCGTGGCCAA GAAGCACCCT AAGACCTGGG TGCACTACAT CGCCGCCGAG
         GAGGAGGACT GGGACTACGC CCCGCTAGTA CTGGCCCCCG ACGACCGCAG
 1251
         CTACAAGAGC CAGTACCTGA ACAACGGCCC CCAGCGCATC GGCCGCAAGT
 1301
 1351
         ACAAGAAGGT GCGCTTCATG GCCTACACCG ACGAGACTTT CAAGACCCGC
1401 GAGGCCATCC AGCACGAGTC CGGCATCCTC GGCCCCCTGC TGTACGGCGA
1451 GGTGGGCGAC ACCCTGCTGA TCATCTTCAA GAACCAGGCC AGCAGGCCCT
1501 ACAACATCTA CCCCCACGGC ATCACCGACG TGCGCCCCCT GTACAGCCGC
1551 CGCCTGCCCA AGGGCGTGAA GCACCTGAAG GACTTCCCCA TCCTGCCCGG
 1601
          CGAGATCTTC AAGTACAAGT GGACCGTGAC CGTGGAGGAC GGCCCCACCA
1651 AGAGCGACCC CCGCTGCCTG ACCCGCTACT ACAGCAGCTT CGTGAACATG
1701 GAGCGCGACC TGGCCTCCGG ACTGATCGGC CCCCTGCTGA TCTGCTACAA
1751 GGAGAGCGTG GACCAGCGCG GCAACCAGAT CATGAGCGAC AAGCGCAACG
         TGATCCTGTT CAGCGTGTTC GACGAGAACC GCAGCTGGTA TCTGACCGAG
        AACATCCAGC GCTTCCTGCC CAACCCCGCT GGCGTGCAGC TGGAAGATCC
CGAGTTCCAG GCCAGCAACA TCATGCACAG CATCAACGGC TACGTGTTCG
ACAGCCTGCA GCTGAGCGTG TGCCTGCATG AGGTGGCCTA CTGGTACATC
1851
1901
1951
2001
         CTGAGCATCG GCGCCCAGAC CGACTTCCTG AGCGTGTTCT TCTCCGGGTA
2051
         TACCTTCAAG CACAAGATGG TGTACGAGGA CACCCTGACC CTGTTCCCCT
         TCTCCGGCGA GACTGTGTTC ATGTCTATGG AGAACCCCGG CCTGTGGATT
CTGGGCTGCC ACAACAGCGA CTTCCGCAAC CGCGGCATGA CTGCCCTGCT
2101
2151
2201
         GAAAGTCTCC AGCTGCGACA AGAACACCGG CGACTACTAC GAGGACAGCT
2251
         ACGAGGACAT CTCCGCCTAC CTGCTGTCCA AGAACAACGC CATCGAGCCC
         CGCTCCTTCT CCCAAAACTC CCGCCACCCC AGCACGCGTC AGAAGCAGTT CAACGCCACC CCCCCGTGC TGAAGCGCCA CCAGCGCGAG ATCACCGCA CCACCCTGCA AAGCGACCAG GAGGAGATCG ACTACGACGA CACCATCAGC
2301
2351
2401
2451
         GTGGAGATGA AGAAGGAGGA CTTCGACATC TACGACGAGG ACGAGAACCA
        GAGCCCCGC TCCTTCCAAA AGAAAACCCG CCACTACTTC ATCGCCGCCG
TGGAGCGCCT GTGGGACTAC GGCATGAGCA GCAGCCCCCA CGTCCTGCGC
AACCGCGCCC AGAGCGGCAG CGTGCCCCAG TTCAAGAAGG TGGTGTTCCA
2501
2551
2601
2651
        GGAGTTCACC GACGGCAGCT TCACCCAGCC CCTGTACCGC GGCGAGCTGA
2701
        ACGAGCACCT GGGCCTGCTC GGCCCCTACA TCCGCGCCGA GGTGGAGGAC
```

Fig. 13

(1 f 2)

				•	4
2751	AACATCATGO			TCCCGGCCCT	ACTCCTTCTA
2801				GCGCCAGGGC	GCCGAGCCCC
2851	GCAAGAACTI				
2901	CAGCACCACA			TTCGACTGCA	AGGCCTGGGC
2951	CTACTTCAGC			CGTGCACAGC	GGCCTGATCG
3001	GCCCCCTGCT				CCACGGGAGG
3051	CAGGTGACTG		TGCCCTGTTC	TTCACCATCT	
3101	TAAGAGCTGG	12112222412	AGAACATGGA	GCGCAACTGC	CGCGCCCCCT
3151	GCAACATCCA		CCCACCTTCA	AGGAGAACTA	
3201	GCCATCAACG		GGACACCCTG	CCCGGCCTGG	TGATGGCCCA
3251	GGACCAGCGC		ACCTGCTGTC	TATGGGCAGC	
3301	TCCACAGCAT		GGCCACGTTT	TCACCGTGCG	CAAGAAGGAG
3351	GAGTACAAGA	TGGCCCTGTA	CAACCTGTAC		TCGAGACTGT
3401	GGAGATGCTG	CCCAGCAAGG	CCGGGATCTG	GCGCGTGGAG	TGCCTGATCG
3451	GCGAGCACCT	GCACGCCGGC	ATGAGCACCC	TGTTCCTGGT	GTACAGCAAC
3501	AAGTGCCAGA	CCCCCTGGG	CATGGCCAGC	GGCCACATCC	GCGACTTCCA
3551	GATCACCGCC	AGCGGCCAGT	ACGGCCAGTG	GGCTCCCAAG	CTGGCCGGCC
3601	TGCACTACAG	CGGCAGCATC	AACGCCTGGT	CGACCAAGGA	GCCCTTCTCC
3651	TGGATCAAGG	TGGACCTGCT	GGCCCCCATG	ATCATCCACG	GCATCAAGAC
3701	CCAGGGCGCC	CGCCAGAAGT	TCAGCAGCCT	GTACATCAGC	CAGTTCATCA
3751	TCATGTACTC	TCTAGACGGC	AAGAAGTGGC	AGACCTACCG	CGGCAACAGC
3801	ACCGGCACCC	TGATGGTGTT	CTTCGGCAAC	GTGGACAGCA	
3851	GCACAACATC		CCATCATCGC	CCGCTACATC	CGCCTGCACC
3901	CCACCCACTA		AGCACCCTGC	GCATGGAGCT.	GATGGGCTGC
3951	GACCTGAACA	GCTGCAGCAT	GCCCCTGGGC	ATGGAGAGCA	AGGCCATCAG
4001	CGACGCCCAG	ATCACCGCCT	CCAGCTACTT	CACCAACATG	TTCGCCACCT
4051	GGAGCCCCAG	CAAGGCCCGC	CTGCACCTGC	AGGGCCGCAG	CAACGCCTGG
4101	CGCCCCAGG	TGAACAACCC	CAAGGAGTGG	CTGCAGGTGG	ACTTCCAGAA
4151	AACCATGAAG	GTGACTGGCG	TGACCACCCA	GGGCGTCAAG	AGCCTGCTGA
4201	CCAGCATGTA	CGTGAAGGAG	TTCCTGATCA	GCAGCAGCCA	GGACGGCCAC
4251	CAGTGGACCC	TGTTCTTCCA	AAACGGCAAG	GTGAAGGTGT	TCCAGGGCAA
4301	CCAGGACAGC	TTCACACCGG	TCGTGAACAG	CCTGGACCCC	CCCCTGCTGA
4351	CCCGCTACCT		CCCCAGAGCT	GGGTGCACCA	GATCGCCCTG
4401	CGCATGGAGG	TGCTGGGCTG	CGAGGCCCAG		GAAGCGGCCG
4451	C i				

Fig. 13

(2 of 2)

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Minimum	documentation searched (classification system follow	red by classification symbols)	
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BIOSIS,	data base consulted during the international search (r EMBASE, MEDLINE, DERWENT		e, search terms used)
	rms: gene?, dna?, nucleic acid?, deoxyribonucleic?,	synthe?, prefer? non-prefer? codon?	
C. DOC	UMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.
X	WO 96/09378 A (THE GENERAL H 28 March 1996, abstract, page 1, line lines 25-32, page 17, lines 27-39 and	20-page 4, line 26, page 15,	1-28
A	SEETHARAM et al. Mistranslation Espression of the Protein in Escherichin Containing Low Frequency Codons. Comm. 30 August 1988. Vol. 155.	1-28	
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(54) Title: HIGH LEVEL EXPRESSION OF PROTEINS

(57) Abstract

The invention features a synthetic gene encoding a protein normally expressed in a mammalian cell wherein at least one non-preferred or less preferred codon in the natural gene encoding the protein has been replaced by a preferred codon encoding the same amino acid.

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HIGH LEVEL EXPRESSION OF PROTEINS

Field of the Invention

The invention concerns genes and methods for expressing eukaryotic and viral proteins at high levels in eukaryotic cells.

Background of the Invention

Expression of eukaryotic gene products in prokaryotes is sometimes limited by the presence of codons that are infrequently used in *E. coli*.

Expression of such genes can be enhanced by systematic substitution of the endogenous codons with codons over represented in highly expressed prokaryotic genes (Robinson et al., Nucleic Acids Res. 12:6663, 1984). It is commonly supposed that rare codons cause pausing of the ribosome, which leads to a failure to complete the nascent polypeptide chain and a uncoupling of transcription and translation. Pausing of the ribosome is thought to lead to exposure of the 3' end of the mRNA to cellular ribonucleases.

Summary of the Invention

The invention features a synthetic gene encoding a protein normally expressed in a mammalian cell or other eukaryotic cell wherein at least one non-preferred or less preferred codon in the natural gene encoding the protein has been replaced by a preferred codon encoding the same amino acid.

Preferred codons are: Ala (gcc); Arg (cgc); Asn (aac); Asp (gac) Cys (tgc); Gln (cag); Gly (ggc); His (cac); Ile (atc); Leu (ctg); Lys (aag); Pro (ccc); Phe (ttc); Ser (agc); Thr (acc); Tyr (tac); and Val (gtg). Less preferred codons are: Gly (ggg); Ile (att); Leu (ctc); Ser (tcc); Val (gtc); and Arg (agg). All codons which do not fit the description of preferred codons or less preferred codons are non-preferred codons. In general, the degree of preference of a particular codon is indicated by the prevalence of the codon in highly expressed human genes as indicated in Table 1 under the heading "High." For example,

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"atc" represents 77% of the Ile codons in highly expressed mammalian genes and is the preferred Ile codon; "att" represents 18% of the Ile codons in highly expressed mammalian genes and is the less preferred Ile codon. The sequence "ata" represents only 5% of the Ile codons in highly expressed human genes as is a non-preferred Ile codon. Replacing a codon with another codon that is more prevalent in highly expressed human genes will generally increase expression of the gene in mammalian cells. Accordingly, the invention includes replacing a less preferred codon with a preferred codon as well as replacing a non-preferred codon with a preferred or less preferred codon.

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By "protein normally expressed in a mammalian cell" is meant a protein which is expressed in mammalian under natural conditions. The term includes genes in the mammalian genome such as those encoding Factor VIII, Factor IX, interleukins, and other proteins. The term also includes genes which are expressed in a mammalian cell under disease conditions such as oncogenes as well as genes which are encoded by a virus (including a retrovirus) which are expressed in mammalian cells post-infection. By "protein normally expressed in a eukaryotic cell" is meant a protein which is expressed in a eukaryote under natural conditions. The term also includes genes which are expressed in a mammalian cell under disease conditions.

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In preferred embodiments, the synthetic gene is capable of expressing the mammalian or eukaryotic protein at a level which is at least 110%, 150%, 200%, 500%, 1,000%, 5,000% or even 10,000% of that expressed by the "natural" (or "native") gene in an *in vitro* mammalian cell culture system under identical conditions (i.e., same cell type, same culture conditions, same expression vector).

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Suitable cell culture systems for measuring expression of the synthetic gene and corresponding natural gene are described below. Other

suitable expression systems employing mammalian cells are well known to those skilled in the art and are described in, for example, the standard molecular biology reference works noted below. Vectors suitable for expressing the synthetic and natural genes are described below and in the standard reference works described below. By "expression" is meant protein expression. Expression can be measured using an antibody specific for the protein of interest. Such antibodies and measurement techniques are well known to those skilled in the art. By "natural gene" and "native gene" is meant the gene sequence (including naturally occurring allelic variants) which naturally encodes the protein, i.e., the native or natural coding sequence.

In other preferred embodiments at least 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, or 90% of the codons in the natural gene are non-preferred codons.

In other preferred embodiments at least 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, or 90% of the non-preferred codons in the natural gene are replaced with preferred codons or less preferred codons.

In other preferred embodiments at least 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, or 90% of the non-preferred codons in the natural gene are replaced with preferred codons.

In a preferred embodiment the protein is a retroviral protein. In a more preferred embodiment the protein is a lentiviral protein. In an even more preferred embodiment the protein is an HIV protein. In other preferred embodiments the protein is gag, pol, env, gp120, or gp160. In other preferred embodiments the protein is a human protein. In more preferred embodiments, the protein is human Factor VIII and the protein in B region deleted human Factor VIII. In another preferred embodiment the protein is green flourescent protein.

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In various preferred embodiments at least 30%, 40%, 50%, 60%, 70%, 80%, 90%, and 95% of the codons in the synthetic gene are preferred or less preferred codons.

The invention also features an expression vector comprising the synthetic gene.

In another aspect the invention features a cell harboring the synthetic gene. In various preferred embodiments the cell is a prokaryotic cell and the cell is a mammalian cell.

In preferred embodiments the synthetic gene includes fewer than 50, fewer than 40, fewer than 30, fewer than 20, fewer than 10, fewer than 5, or no "cg" sequences.

The invention also features a method for preparing a synthetic gene encoding a protein normally expressed by a mammalian cell or other eukaryotic cell. The method includes identifying non-preferred and less-preferred codons in the natural gene encoding the protein and replacing one or more of the non-preferred and less-preferred codons with a preferred codon encoding the same amino acid as the replaced codon.

Under some circumstances (e.g., to permit introduction of a restriction site) it may be desirable to replace a non-preferred codon with a less preferred codon rather than a preferred codon.

It is not necessary to replace all less preferred or non-preferred codons with preferred codons. Increased expression can be accomplished even with partial replacement of less preferred or non-preferred codons with preferred codons. Under some circumstances it may be desirable to only partially replace non-preferred codons with preferred or less preferred codons in order to obtain an intermediate level of expression.

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In other preferred embodiments the invention features vectors (including expression vectors) comprising one or more the synthetic genes.

By "vector" is meant a DNA molecule, derived, e.g., from a plasmid, bacteriophage, or mammalian or insect virus, into which fragments of DNA may be inserted or cloned. A vector will contain one or more unique restriction sites and may be capable of autonomous replication in a defined host or vehicle organism such that the cloned sequence is reproducible. Thus, by "expression vector" is meant any autonomous element capable of directing the synthesis of a protein. Such DNA expression vectors include mammalian plasmids and viruses.

The invention also features synthetic gene fragments which encode a desired portion of the protein. Such synthetic gene fragments are similar to the synthetic genes of the invention except that they encode only a portion of the protein. Such gene fragments preferably encode at least 50, 100, 150, or 500 contiguous amino acids of the protein.

In constructing the synthetic genes of the invention it may be desirable to avoid CpG sequences as these sequences may cause gene silencing. Thus, in a preferred embodiment the coding region of the synthetic gene does not include the sequence "cg."

The codon bias present in the HIV gp120 env gene is also present in the gag and pol genes. Thus, replacement of a portion of the non-preferred and less preferred codons found in these genes with preferred codons should produce a gene capable of higher level expression. A large fraction of the codons in the human genes encoding Factor VIII and Factor IX are non-preferred codons or less preferred codons. Replacement of a portion of these codons with preferred codons should yield genes capable of higher level expression in mammalian cell culture.

The synthetic genes of the invention can be introduced into the cells of a living organism. For example, vectors (viral or non-viral) can be used to introduce a synthetic gene into cells of a living organism for gene therapy.

Conversely, it may be desirable to replace preferred codons in a naturally occurring gene with less-preferred codons as a means of lowering expression.

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Standard reference works describing the general principles of recombinant DNA technology include Watson et al., Molecular Biology of the Gene, Volumes I and II, the Benjamin/Cummings Publishing Company, Inc.,

publisher, Menlo Park, CA (1987); Darnell et al., Molecular Cell Biology,
Scientific American Books, Inc., Publisher, New York, N.Y. (1986); Old et al.,
Principles of Gene Manipulation: An Introduction to Genetic Engineering, 2d edition, University of California Press, publisher, Berkeley, CA (1981);
Maniatis et al., Molecular Cloning: A Laboratory Manual, 2nd Ed. Cold Spring
Harbor Laboratory, publisher, Cold Spring Harbor, NY (1989); and Current Protocols in Molecular Biology, Ausubel et al., Wiley Press, New York, NY (1992).

By "transformed cell" is meant a cell into which (or into an ancestor of which) has been introduced, by means of recombinant DNA techniques, a selected DNA molecule, e.g., a synthetic gene.

By "positioned for expression" is meant that a DNA molecule, e.g., a synthetic gene, is positioned adjacent to a DNA sequence which directs transcription and translation of the sequence (i.e., facilitates the production of the protein encoded by the synthetic gene.

Description of the Drawings

Figure 1 depicts the sequence of the synthetic gp120 and a synthetic gp160 gene in which codons have been replaced by those found in highly expressed human genes.

Figure 2 is a schematic drawing of the synthetic gp120 (HIV-1 MN) gene. The shaded portions marked v1 to v5 indicate hypervariable regions. The filled box indicates the CD4 binding site. A limited number of the unique restriction sites ares shown: H (Hind3), Nh (Nhe1), P (Pst1), Na (Nae1), M (Mlu1), R (EcoR1), A (Age1) and No (Not1). The chemically synthesized DNA fragments which served as PCR templates are shown below the gp120 sequence, along with the locations of the primers used for their amplification.

Figure 3 is a photograph of the results of transient transfection assays used to measure gp120 expression. Gel electrophoresis of immunoprecipitated supernatants of 293T cells transfected with plasmids expressing gp120 encoded by the IIIB isolate of HIV-1 (gp120IIIb), by the MN isolate of HIV-1 (gp120mn), by the MN isolate of HIV-1 modified by substitution of the endogenous leader peptide with that of the CD5 antigen (gp120mnCD5L), or by the chemically synthesized gene encoding the MN variant of HIV-1 with the human CD5Leader (syngp120mn). Supernatants were harvested following a 12 hour labeling period 60 hours post-transfection and immunoprecipitated with CD4:IgG1 fusion protein and protein A sepharose.

Figure 4 is a graph depicting the results of ELISA assays used to measure protein levels in supernatants of transiently transfected 293T cells. Supernatants of 293T cells transfected with plasmids expressing gp120 encoded by the IIIB isolate of HIV-1 (gp120 IIIb), by the MN isolate of HIV-1 (gp120mn), by the MN isolate of HIV-1 modified by substitution of the endogenous leader peptide with that of CD5 antigen (gp120mn CD5L), or by

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the chemically synthesized gene encoding the MN variant of HIV-1 with human CDS leader (syngp120mn) were harvested after 4 days and tested in a gp120/CD4 ELISA. The level of gp120 is expressed in ng/ml.

Figure 5A is a photograph of a gel illustrating the results of a immunoprecipitation assay used to measure expression of the native and synthetic gp120 in the presence of rev in trans and the RRE in cis. In this experiment 293T cells were transiently transfected by calcium phosphate coprecipitation of 10 μ g of plasmid expressing: (A) the synthetic gp120MN sequence and RRE in cis, (B) the gp120 portion of HIV-1 IIIB, (C) the gp120 portion of HIV-1 IIIB and RRE in cis, all in the presence or absence of rev expression. The RRE constructs gp120IIIbRRE and syngp120mnRRE were generated using an Eag1/Hpa1 RRE fragment cloned by PCR from a HIV-1 HXB2 proviral clone. Each gp120 expression plasmid was cotransfected with 10 μ g of either pCMVrev or CDM7 plasmid DNA. Supernatants were harvested 60 hours post transfection, immunoprecipitated with CD4:IgG fusion protein and protein A agarose, and run on a 7% reducing SDS-PAGE. The gel exposure time was extended to allow the induction of gp120IIIbrre by rev to be demonstrated.

z Figure 5B is a shorter exposure of a similar experiment in which syngp120mnrre was cotransfected with or without pCMVrev.

Figure 5C is a schematic diagram of the constructs used in Figure 5A.

Figure 6 is a comparison of the sequence of the wild-type ratTHY-1 gene (wt) and a synthetic ratTHY-1 gene (env) constructed by chemical synthesis and having the most prevalent codons found in the HIV-1 env gene.

Figure 7 is a schematic diagram of the synthetic ratTHY-1 gene. The solid black box denotes the signal peptide. The shaded box denotes the

sequences in the precursor which direct the attachment of a phophatidylinositol glycan anchor. Unique restriction sites used for assembly of the THY-1 constructs are marked H (Hind3), M (Mlu1), S (Sac1) and No (Not1). The position of the synthetic oligonucleotides employed in the construction are shown at the bottom of the figure.

Figure 8 is a graph depicting the results of flow cytometry analysis. In this experiment 293T cells transiently transfected with either a wild-type ratTHY-1 expression plasmid (thick line), ratTHY-1 with envelope codons expression plasmid (thin line), or vector only (dotted line) by calcium phosphate co-precipitation. Cells were stained with anti-ratTHY-1 monoclonal antibody OX7 followed by a polyclonal FITC-conjugated anti-mouse IgG antibody 3 days after transfection.

Figure 9A is a photograph of a gel illustrating the results of immunoprecipitation analysis of supernatants of human 293T cells transfected with either syngp120mn (A) or a construct syngp120mn.rTHY-1env which has the rTHY-1env gene in the 3' untranslated region of the syngp120mn gene (B). The syngp120mn.rTHY-1env construct was generated by inserting a Not1 adapter into the blunted Hind3 site of the rTHY-1env plasmid. Subsequently, a 0.5 kb Not1 fragment containing the rTHY-1env gene was cloned into the Not1 site of the syngp120mn plasmid and tested for correct orientation. Supernatants of ³⁵S labeled cells were harvested 72 hours post transfection, precipitated with CD4:IgG fusion protein and protein A agarose, and run on a 7% reducing SDS-PAGE.

Figure 9B is a schematic diagram of the constructs used in the experiment depicted in Figure 9A.

Figure 10A is a photograph of COS cells transfected with vector only showing no GFP fluorescence.

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Figure 10B is a photograph of COS cells transfected with a CDM7 expression plasmid encoding native GFP engineered to include a consensus translational initiation sequence.

Figure 10C is a photograph of COS cells transfected with an expression plasmid having the same flanking sequences and initiation consensus as in Figure 10B, but bearing a codon optimized gene sequence.

Figure 10D is a photograph of COS cells transfected with an expression plasmid as in Figure 10C, but bearing a Thr at residue 65 in place of Ser.

Figure 11 depicts the sequence of a synthetic gene encoding green flourescent proteins (SEQ ID NO:40).

Figure 12 depicts the sequence of a native human Factor VIII gene lacking the central B domain (amino acids 760-1639, inclusive) (SEQ ID NO:41).

Figure 13 depicts the sequence of a synthetic human Factor VIII gene lacking the central B domain (amino acids 760-1639, inclusive) (SEQ ID NO:42).

Description of the Preferred Embodiments

EXAMPLE 1

20 Construction of a Synthetic gp120 Gene Having Codons Found in Highly Expressed Human Genes

A codon frequency table for the envelope precursor of the LAV subtype of HIV-1 was generated using software developed by the University of Wisconsin Genetics Computer Group. The results of that tabulation are contrasted in Table 1 with the pattern of codon usage by a collection of highly expressed human genes. For any amino acid encoded by degenerate codons,

the most favored codon of the highly expressed genes is different from the most favored codon of the HIV envelope precursor. Moreover a simple rule describes the pattern of favored envelope codons wherever it applies: preferred codons maximize the number of

adenine residues in the viral RNA. In all cases but one this means that the codon in which the third position is A is the most frequently used. In the special case of serine, three codons equally contribute one A residue to the mRNA; together these three comprise 85% of the serine codons actually used in envelope transcripts. A particularly striking example of the A bias is found in the codon choice for arginine, in which the AGA triplet comprises 88% of the arginine codons. In addition to the preponderance of A residues, a marked preference is seen for uridine among degenerate codons whose third residue must be a pyrimidine. Finally, the inconsistencies among the less frequently used variants can be accounted for by the observation that the dinucleotide CpG is under represented; thus the third position is less likely to be G whenever the second position is C, as in the codons for alanine, proline, serine and threonine; and the CGX triplets for arginine are hardly used at all.

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TABLE 1:	,	Codon Frequency in the HIV-1 IIIb env gene and in high	ly
		expressed human genes.	
		OND, ODOG MANAGEMENT	1.

		1	1		' '	• •	.,	•	L =	_ '
			High	Env		•		,	High	Env
	Ala						<u>Cys</u> '		1	
5	GC -	$\mathbf{C}^{'}$	53	27	•	1 .	TG	\cdot C	68 .	16
		\mathbf{T}_{1}	17	18	1		, '	T	32	84
		\mathbf{A}	. 13 "	50 '				, 1	1	1
	. :	G	17	5	• •		<u>Gln</u>	130		٠.
				. ' 1	•		CA',	Α	12	55
10	Arg					•		, G	88	45 ,
•	CG	C	, 37	0			1 1			
		T	7	4	•		<u>Glu</u>			
		${f A}$,	6	0		•	GA	A	25	67
		G	21	. 0	**			\mathbf{G}^{T}	['] 75	33
15	AG	A	10	88				1	•	
		G [']	18	8		i	<u>Gly</u>			
		_	1			. '	GG	C	50	6
	<u>Asn</u>			t		ľ		T	12	13
	AA	С	78	30				Α	14	53
20		T	. 22	70				G	24	28
	٠	. -								
	Asp					1	<u>His</u>			
	GA	\mathbf{C}	75	33			CA	C	79	25
		T	25	67				T	21	75
	,	_		-						
							<u>Ile</u>	,		•
25						٠.	$\overline{\mathbf{A}}\mathbf{T}$	C	77	25
			+					T	18	31
							•	Α	5 .	44
			1						1	
	Leu						Ser			
	CT	С	26	10			TC	C	28	8
30		T	5	7				T	13	8
~ ~		Ā	3	17				Α	5	22
		G	58	17				G	9	0
	TT	Ā	2	30			AG	C	34	22
		G	6	20				T	10	41

ł	<u>Lys</u>		•			•	Thr	· _		
	AA	\mathbf{A}	18	68	٠,		AC	, C	57	20
	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	G	82	32	1	. 4		\mathbf{T}_{\cdot}	14	22
								A	14	51
5								G	15	7
,	. 1				•					
	Pro .					1	Tyr			
	· CC	C	48	27	ı	' '	TA	C	74	8
		T	19	14 '				T	· 26	92
		\mathbf{A}	16	55				.,		
10	1	G	17	5						
	1	. ,		•						
	Phe						<u>Val</u>		,	
	TT	C	80	26			GT	C	25	12
		T	20	74				T	7	9 ,
								Α	5	62
15	1							G	64	18

Codon frequency was calculated using the GCG program established the University of Wisconsin Genetics Computer Group. Numbers represent the percentage of cases in which the particular codon is used. Codon usage frequencies of envelope genes of other HIV-1 virus isolates are comparable and show a similar bias.

In order to produce a gp120 gene capable of high level expression in mammalian cells, a synthetic gene encoding the gp120 segment of HIV-1 was constructed (syngp120mn), based on the sequence of the most common North American subtype, HIV-1 MN (Shaw et al., Science 226:1165, 1984; Gallo et al., Nature 321:119, 1986). In this synthetic gp120 gene nearly all of the native codons have been systematically replaced with codons most frequently used in highly expressed human genes (Figure 1). This synthetic gene was assembled from chemically synthesized oligonucleotides of 150 to 200 bases in length. If oligonucleotides exceeding 120 to 150 bases are chemically synthesized, the

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percentage of full-length product can be low, and the vast excess of material consists of shorter oligonucleotides. Since these shorter fragments inhibit cloning and PCR procedures, it can be very difficult to use oligonucleotides exceeding a certain length. In order to use crude synthesis material without prior purification, single-stranded oligonucleotide pools were PCR amplified before cloning. PCR products were purified in agarose gels and used as templates in the next PCR step. Two adjacent fragments could be co-amplified because of overlapping sequences at the end of either fragment. These fragments, which were between 350 and 400 bp in size, were subcloned into a pCDM7-derived plasmid containing the leader sequence of the CD5 surface molecule followed by a Nhe1/Pst1/Mlu1/EcoR1/BamH1 polylinker. Each of the restriction enzymes in this polylinker represents a site that is present at either the 5' or 3' end of the PCR-generated fragments. Thus, by sequential subcloning of each of the 4 long fragments, the whole gp120 gene was assembled. For each fragment three to six different clones were subcloned and sequenced prior to assembly. A schematic drawing of the method used to construct the synthetic gp120 is shown in Figure 2. The sequence of the synthetic gp120 gene (and a synthetic gp160 gene created using the same approach) is presented in Figure 1.

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The mutation rate was considerable. The most commonly found mutations were short (1 nucleotide) and long (up to 30 nucleotides) deletions.

In some cases it was necessary to exchange parts with either synthetic adapters or pieces from other subclones without mutation in that particular region.

Some deviations from strict adherence to optimized codon usage were made to accommodate the introduction of restriction sites into the resulting gene to facilitate the replacement of various segments (Figure 2). These unique restriction sites were introduced into the gene at approximately 100 bp

intervals. The native HIV leader sequence was exchanged with the highly efficient leader peptide of the human CD5 antigen to facilitate secretion (Aruffo et al., Cell 61:1303, 1990) The plasmid used for construction is a derivative of the mammalian expression vector pCDM7 transcribing the inserted gene under the control of a strong human CMV immediate early promoter.

To compare the wild-type and synthetic gp120 coding sequences, the synthetic gp120 coding sequence was inserted into a mammalian expression vector and tested in transient transfection assays. Several different native gp120 genes were used as controls to exclude variations in expression levels between different virus isolates and artifacts induced by distinct leader sequences. The gp120 HIV IIIb construct used as control was generated by PCR using a Sal1/Xho1 HIV-1 HXB2 envelope fragment as template. To exclude PCR induced mutations, a Kpn1/Ear1 fragment containing approximately 1.2 kb of the gene was exchanged with the respective sequence from the proviral clone. The wild-type gp120mn constructs used as controls were cloned by PCR from HIV-1 MN infected C8166 cells (AIDS Repository, Rockville, MD) and expressed gp120 either with a native envelope or a CD5 leader sequence. Since proviral clones were not available in this case, two clones of each construct were tested to avoid PCR artifacts. To determine the amount of secreted gp120 semi-quantitatively supernatants of 293T cells transiently transfected by calcium phosphate co-precipitation were immunoprecipitated with soluble CD4:immunoglobulin fusion protein and protein A sepharose.

The results of this analysis (Figure 3) show that the synthetic gene product is expressed at a very high level compared to that of the native gp120 controls. The molecular weight of the synthetic gp120 gene was comparable to

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control proteins (Figure 3) and appeared to be in the range of 100 to 110 kd. The slightly faster migration can be explained by the fact that in some tumor cell lines, e.g., 293T, glycosylation is either not complete or altered to some extent.

To compare expression more accurately gp120 protein levels were quantitated using a gp120 ELISA with CD4 in the demobilized phase. This analysis shows (Figure 4) that ELISA data were comparable to the immunoprecipitation data, with a gp120 concentration of approximately 125 ng/ml for the synthetic gp120 gene, and less than the background cutoff (5 ng/ml) for all the native gp120 genes. Thus, expression of the synthetic gp120 gene appears to be at least one order of magnitude higher than wild-type gp120 genes. In the experiment shown the increase was at least 25 fold.

The Role of rev in gp120 Expression

Since rev appears to exert its effect at several steps in the expression of a viral transcript, the possible role of non-translational effects in the improved expression of the synthetic gp120 gene was tested. First, to rule out the possibility that negative signals elements conferring either increased mRNA degradation or nucleic retention were eliminated by changing the nucleotide sequence, cytoplasmic mRNA levels were tested. Cytoplasmic RNA was prepared by NP40 lysis of transiently transfected 293T cells and subsequent elimination of the nuclei by centrifugation. Cytoplasmic RNA was subsequently prepared from lysates by multiple phenol extractions and precipitation, spotted on nitrocellulose using a slot blot apparatus, and finally hybridized with an envelope-specific probe.

Briefly, cytoplasmic mRNA 293 cells transfected with CDM&, gp120 IIIB, or syngp120 was isolated 36 hours post transfection. Cytoplasmic RNA of Hela cells infected with wild-type vaccinia virus or recombinant virus

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expressing gp120 IIIb or the synthetic gp120 gene was under the control of the 7.5 promoter was isolated 16 hours post infection. Equal amounts were spotted on nitrocellulose using a slot blot device and hybridized with randomly labeled 1.5 kb gp120IIIb and syngp120 fragments or human beta-actin. RNA expression levels were quantitated by scanning the hybridized membranes with a phospoimager. The procedures used are described in greater detail below.

This experiment demonstrated that there was no significant difference in the mRNA levels of cells transfected with either the native or synthetic gp120 gene. In fact, in some experiments cytoplasmic mRNA level of the synthetic gp120 gene was even lower than that of the native gp120 gene.

These data were confirmed by measuring expression from recombinant vaccinia viruses. Human 293 cells or Hela cells were infected with vaccinia virus expressing wild-type gp120 IIIb or syngp120mn at a multiplicity of infection of at least 10. Supernatants were harvested 24 hours post infection and immunoprecipitated with CD4:immunoglobin fusion protein and protein A sepharose. The procedures used in this experiment are described in greater detail below.

This experiment showed that the increased expression of the synthetic gene was still observed when the endogenous gene product and the synthetic gene product were expressed from vaccinia virus recombinants under the control of the strong mixed early and late 7.5k promoter. Because vaccinia virus mRNAs are transcribed and translated in the cytoplasm, increased expression of the synthetic envelope gene in this experiment cannot be attributed to improved export from the nucleus. This experiment was repeated in two additional human cell types, the kidney cancer cell line 293 and HeLa cells. As with transfected 293T cells, mRNA levels were similar in 293 cells infected with either recombinant vaccinia virus.

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Codon Usage in Lentivirus

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9812207A1 IA:

Because it appears that codon usage has a significant impact on expression in mammalian cells, the codon frequency in the envelope genes of other retroviruses was examined. This study found no clear pattern of codon preference between retroviruses in general. However, if viruses from the lentivirus genus, to which HIV-1 belongs to, were analyzed separately, codon usage bias almost identical to that of HIV-1 was found. A codon frequency table from the envelope glycoproteins of a variety of (predominantly type C) retroviruses excluding the lentiviruses was prepared, and compared a codon frequency table created from the envelope sequences of four lentiviruses not closely related to HIV-1 (caprine arthritis encephalitis virus, equine infectious anemia virus, feline immunodeficiency virus, and visna virus) (Table 2). The codon usage pattern for lentiviruses is strikingly similar to that of HIV-1, in all cases but one, the preferred codon for HIV-1 is the same as the preferred codon for the other lentiviruses. The exception is proline, which is encoded by CCT in 41% of non-HIV lentiviral envelope residues, and by CCA in 40% of residues, a situation which clearly also reflects a significant preference for the triplet ending in A. The pattern of codon usage by the non-lentiviral envelope proteins does not show a similar predominance of A residues, and is also not as skewed toward third position C and G residues as is the codon usage for the highly expressed human genes. In general non-lentiviral retroviruses appear to exploit the different codons more equally, a pattern they share with less highly expressed human genes.

TABLE 2: Codon frequency in the envelope gene of lentiviruses (lenti) and non-lentiviral retroviruses (other)

		Ó	Other Lenti							
	<u>Ala</u>			. '			Cys	1	i	
5	GC.	C	45	13				. C	53	21
•	G G .		26		ı	1	,	Ť	47	79
		T _r	20			٠.	· . •	•		. ,
		G	9				<u>Gln</u>	14		
		J	<i>.</i>	3			CA		52	69
1.0	A				,				48	
10	Arg	~	1.4	٠,			1 4	G	40	31
	CG	C	, 14	2 , 3 ,			C 1			
		T	6	13,	·		<u>Glu</u>	,		
		Α,	16	5			GA	A	57	68
		G	17	3				G	43	32
15	AG	\mathbf{A}	31				, .			
		G	15	26		, 1	<u>Gly</u>			
			1, ,		,		GG	C	21	8
	<u>Asn</u>					. '		T	13	9
	AA	C	49	31				Α	37	56
20		T	51	69				G	29	26
			•							
	<u>Asp</u>						<u>His</u>			
	GĀ	\mathbf{C}	55	33			CA	C	51	38
		Т	51	69				T	49	62
										•
							<u>Ile</u>			
25							AT	С	38	16
			•					T	31	22
								Ā	31	61
			1							01
	<u>Leu</u>						Ser		•	
	CT	C	22	8			TC	С	38	10
30	Cı	T		9			10	T	17	
30		A		16					18	24
								A G		5
	Tr.Tr	G	19	11			A C		6	
	TT	A		41			AG	C	13	20
		G	10	16				T	7	25

					,						
!	1				1 '	•				•	1
` 	Lys		4		٠	•	Thr	4			
	ĀĀ	A	60	63	٠.,		AC	C	44	1.8	
	, •	A G	· 40	37	i			T	27	20	
					•			A	19	• 55	
5	Pro							G	10	8	
	CC	\mathbf{C}	42	14	•						,
	,	T	30	41		t	Tyr				
	. 10	Α	20	40	. '	. '	ΤĄ	C	48	28	
		G	7	5				T	. 52	72	
											t
10	Phe		•			•	<u>Val</u>		•		
	TT	\mathbf{C} . ,	52	25			GT	C	36	9	
		T	48	75				T '	17	10	
•	1						•	A	22	54	
								G	25	27	1
										•	

PCT/US97/16639

Codon frequency was calculated using the GCG program established by the University of Wisconsin Genetics Computer Group. Numbers represent the percentage in which a particular codon is used. Codon usage of non-lentiviral retroviruses was compiled from the envelope precursor sequences of bovine leukemia virus feline leukemia virus, human T-cell leukemia virus type I, human T-cell lymphotropic virus type II, the mink cell focus-forming isolate of murine leukemia virus (MuLV), the Rauscher spleen focus-forming isolate, the 10A1 isolate, the 4070A amphotropic isolate and the myeloproliferative leukemia virus isolate, and from rat leukemia virus, simian sarcoma virus, simian T-cell leukemia virus, leukemogenic retrovirus T1223/B and gibbon ape leukemia virus. The codon frequency tables for the non-HIV, non-SIV lentiviruses were compiled from the envelope precursor sequences for caprine arthritis encephalitis virus, equine infectious anemia virus, feline immunodeficiency virus, and visna virus.

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In addition to the prevalence of codons containing an A, lentiviral codons adhere to the HIV pattern of strong CpG under representation, so that the third position for alanine, proline, serine and threonine triplets is rarely G. The retroviral envelope triplets show a similar, but less pronounced, under representation of CpG. The most obvious difference between lentiviruses and

other retroviruses with respect to CpG prevalence lies in the usage of the CGX variant of arginine triplets, which is reasonably frequently represented among the retroviral envelope coding sequences, but is almost never present among the comparable lentivirus sequences.

5 Differences in rev Dependence Between Native and Synthetic gp120

To examine whether regulation by rev is connected to HIV-1 codon usage, the influence of rev on the expression of both native and synthetic gene was investigated. Since regulation by rev requires the rev-binding site RRE in cis, constructs were made in which this binding site was cloned into the 3' untranslated region-of both the native and the synthetic gene. These plasmids were co-transfected with rev or a control plasmid in trans into 293T cells, and gp120 expression levels in supernatants were measured semiquantitatively by immunoprecipitation. The procedures used in this experiment are described in greater detail below.

As shown in Figure 5A and Figure 5B, rev up regulates the native gp120 gene, but has no effect on the expression of the synthetic gp120 gene. Thus, the action of rev is not apparent on a substrate which lacks the coding sequence of endogenous viral envelope sequences.

Expression of a synthetic ratTHY-1 gene with HIV envelope codons

The above-described experiment suggest that in fact "envelope sequences" have to be present for rev regulation. In order to test this hypothesis, a synthetic version of the gene encoding the small, typically highly expressed cell surface protein, ratTHY-1 antigen, was prepared. The synthetic version of the ratTHY-1 gene was designed to have a codon usage like that of HIV gp120. In designing this synthetic gene AUUUA sequences, which are associated with mRNA instability, were avoided. In addition, two restriction

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sites were introduced to simplify manipulation of the resulting gene (Figure 6). This synthetic gene with the HIV envelope codon usage (rTHY-1env) was generated using three 150 to 170 mer oligonucleotides (Figure 7). In contrast to the syngp120mn gene, PCR products were directly cloned and assembled in pUC12, and subsequently cloned into pCDM7.

Expression levels of native rTHY-1 and rTHY-1 with the HIV envelope codons were quantitated by immunofluorescence of transiently transfected 293T cells. Figure 8 shows that the expression of the native THY-1 gene is almost two orders of magnitude above the background level of the control transfected cells (pCDM7). In contrast, expression of the synthetic ratTHY-1 is substantially lower than that of the native gene (shown by the shift to of the peak towards a lower channel number).

To prove that no negative sequence elements promoting mRNA degradation were inadvertently introduced, a construct was generated in which the rTHY-1env gene was cloned at the 3' end of the synthetic gp120 gene (Figure 9B). In this experiment 293T cells were transfected with either the syngp120mn gene or the syngp120/ratTHY-1 env fusion gene (syngp120mn.rTHY-1env). Expression was measured by immunoprecipitation with CD4:IgG fusion protein and protein A agarose. The procedures used in this experiment are described in greater detail below.

Since the synthetic gp120 gene has an UAG stop codon, rTHY-1env is not translated from this transcript. If negative elements conferring enhanced degradation were present in the sequence, gp120 protein levels expressed from this construct should be decreased in comparison to the syngp120mn construct without rTHY-1env. Figure 9A, shows that the expression of both constructs is similar, indicating that the low expression must be linked to translation.

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Rev-dependent expression of synthetic ratTHY-1 gene with envelope codons

To explore whether rev is able to regulate expression of a ratTHY-1 gene having env codons, a construct was made with a rev-binding site in the 3' end of the rTHY1env open reading frame. To measure rev-responsiveness of the a ratTHY-1env construct having a 3' RRE, human 293T cells were cotransfected ratTHY-1envrre and either CDM7 or pCMVrev. At 60 hours post transfection cells were detached with 1 mM EDTA in PBS and stained with the OX-7 anti rTHY-1 mouse monoclonal antibody and a secondary FITC-conjugated antibody. Fluorescence intensity was measured using a EPICS XL cytofluorometer. These procedures are described in greater detail below.

In repeated experiments, a slight increase of rTHY-1env expression was detected if rev was cotransfected with the rTHY-1env gene. To further increase the sensitivity of the assay system a construct expressing a secreted version of rTHY-1env was generated. This construct should produce more reliable data because the accumulated amount of secreted protein in the supernatant reflects the result of protein production over an extended period, in contrast to surface expressed protein, which appears to more closely reflect the current production rate. A gene capable of expressing a secreted form was prepared by PCR using forward and reverse primers annealing 3' of the endogenous leader sequence and 5' of the sequence motif required for phosphatidylinositol glycan anchorage respectively. The PCR product was cloned into a plasmid which already contained a CD5 leader sequence, thus generating a construct in which the membrane anchor has been deleted and the leader sequence exchanged by a heterologous (and probably more efficient) leader peptide.

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The rev-responsiveness of the secreted form ratTHY-1env was measured by immunoprecipitation of supernatants of human 293T cells cotransfected with a plasmid expressing a secreted form of ratTHY-1env and the RRE sequence in cis (rTHY-1envPI-rre) and either CDM7 or pCMVrev. The rTHY-1envPI-RRE construct was made by PCR using the oligonucleotide: cgcggggctagcgcaaagagtaataagtttaac (SEQ ID NO:38) as a forward primer, the oligonucleotide: cgcggatcccttgtattttgtactaata (SEQ ID NO:39) as reverse primer, and the synthetic rTHY-1env construct as a template. After digestion with Nhe1 and Not1 the PCR fragment was cloned into a plasmid containing CD5 leader and RRE sequences. Supernatants of ³⁵S labeled cells were harvested 72 hours post transfection, precipitated with a mouse monoclonal antibody OX7 against rTHY-1 and anti mouse IgG sepharose, and run on a 12% reducing SDS-PAGE.

In this experiment the induction of rTHY-1env by rev was much more prominent and clear-cut than in the above-described experiment and strongly suggests that rev is able to translationally regulate transcripts that are suppressed by low-usage codons.

Rev-independent expression of a rTHY-lenv:immunoglobulin fusion protein

To test whether low-usage codons must be present throughout the whole coding sequence or whether a short region is sufficient to confer revresponsiveness, a rTHY-lenv:immunoglobulin fusion protein was generated. In this construct the rTHY-lenv gene (without the sequence motif responsible for phosphatidylinositol glycan anchorage) is linked to the human IgG1 hinge, CH2 and CH3 domains. This construct was generated by anchor PCR using primers with Nhe1 and BamHI restriction sites and rTHY-lenv as template. The PCR fragment was cloned into a plasmid containing the leader sequence of

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the CD5 surface molecule and the hinge, CH2 and CH3 parts of human IgG1 immunoglobulin. A Hind3/Eag1 fragment containing the rTHY-1enveg1 insert was subsequently cloned into a pCDM7-derived plasmid with the RRE sequence.

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To measure the response of the rTHY-1env/ immunoglobin fusion gene (rTHY-1enveg1rre) to rev human 293T cells cotransfected with rTHY-1enveg1rre and either pCDM7 or pCMVrev. The rTHY-1enveg1rre construct was made by anchor PCR using forward and reverse primers with Nhe1 and BamH1 restriction sites respectively. The PCR fragment was cloned—into-a-plasmid-containing-a-CD5-leader-and-human IgG1 hinge, CH2 and CH3 domains. Supernatants of ³⁵S labeled cells were harvested 72 hours post transfection, precipitated with a mouse monoclonal antibody OX7 against rTHY-1 and anti mouse IgG sepharose, and run on a 12% reducing SDS-PAGE. The procedures used are described in greater detail below.

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As with the product of the rTHY-1envPI- gene, this rTHY-1env/immunoglobulin fusion protein is secreted into the supernatant. Thus, this gene should be responsive to rev-induction. However, in contrast to rTHY-1envPI-, cotransfection of rev in trans induced no or only a negligible increase of rTHY-1enveg1 expression.

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The expression of rTHY-1:immunoglobulin fusion protein with native rTHY-1 or HIV envelope codons was measured by immunoprecipitation. Briefly, human 293T cells transfected with either rTHY-1enveg1 (env codons) or rTHY-1wteg1 (native codons). The rTHY-1wteg1 construct was generated in manner similar to that used for the rTHY-1enveg1 construct, with the exception that a plasmid containing the native rTHY-1 gene was used as template. Supernatants of ³⁵S labeled cells were harvested 72 hours post transfection, precipitated with a mouse monoclonal antibody OX7 against

rTHY-1 and anti mouse IgG sepharose, and run on a 12% reducing SDS-PAGE. THE procedures used in this experiment are described in greater detail below.

Expression levels of rTHY-1enveg1 were decreased in comparison to a similar construct with wild-type rTHY-1 as the fusion partner, but were still considerably higher than rTHY-1env. Accordingly, both parts of the fusion protein influenced expression levels. The addition of rTHY-1env did not restrict expression to an equal level as seen for rTHY-1env alone. Thus, regulation by rev appears to be ineffective if protein expression is not almost completely suppressed.

Codon preference in HIV-1 envelope genes

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Direct comparison between codon usage frequency of HIV envelope and highly expressed human genes reveals a striking difference for all twenty amino acids. One simple measure of the statistical significance of this codon preference is the finding that among the nine amino acids with two fold codon degeneracy, the favored third residue is A or U in all nine. The probability that all nine of two equiprobable choices will be the same is approximately 0.004, and hence by any conventional measure the third residue choice cannot be considered random. Further evidence of a skewed codon preference is found among the more degenerate codons, where a strong selection for triplets bearing adenine can be seen. This contrasts with the pattern for highly expressed genes, which favor codons bearing C, or less commonly G, in the third position of codons with three or more fold degeneracy.

The systematic exchange of native codons with codons of highly expressed human genes dramatically increased expression of gp120. A quantitative analysis by ELISA showed that expression of the synthetic gene was at least 25 fold higher in comparison to native gp120 after transient

transfection into human 293 cells. The concentration levels in the ELISA experiment shown were rather low. Since an ELISA was used for quantification which is based on gp120 binding to CD4, only native, non-denatured material was detected. This may explain the apparent low expression. Measurement of cytoplasmic mRNA levels demonstrated that the difference in protein expression is due to translational differences and not mRNA stability.

Retroviruses in general do not show a similar preference towards A and T as found for HIV. But if this family was divided into two subgroups, lentiviruses and non-lentiviral-retroviruses, a-similar preference to A and, less frequently, T, was detected at the third codon position for lentiviruses. Thus, the availing evidence suggests that lentiviruses retain a characteristic pattern of envelope codons not because of an inherent advantage to the reverse transcription or replication of such residues, but rather for some reason peculiar to the physiology of that class of viruses. The major difference between lentiviruses and non-complex retroviruses are additional regulatory and nonessentially accessory genes in lentiviruses, as already mentioned. Thus, one simple explanation for the restriction of envelope expression might be that an important regulatory mechanism of one of these additional molecules is based on it. In fact, it is known that one of these proteins, rev, which most likely has homologues in all lentiviruses. Thus codon usage in viral mRNA is used to create a class of transcripts which is susceptible to the stimulatory action of rev. This hypothesis was proved using a similar strategy as above, but this time codon usage was changed into the inverse direction. Codon usage of a highly expressed cellular gene was substituted with the most frequently used codons in the HIV envelope. As assumed, expression levels were considerably lower in comparison to the native molecule, almost two orders of magnitude when

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analyzed by immunofluorescence of the surface expressed molecule. If rev was coexpressed in trans and a RRE element was present in cis only a slight induction was found for the surface molecule. However, if THY-1 was expressed as a secreted molecule, the induction by rev was much more prominent, supporting the above hypothesis. This can probably be explained by accumulation of secreted protein in the supernatant, which considerably amplifies the rev effect. If rev only induces a minor increase for surface molecules in general, induction of HIV envelope by rev cannot have the purpose of an increased surface abundance, but rather of an increased intracellular gp160 level. It is completely unclear at the moment why this should be the case.

To test whether small subtotal elements of a gene are sufficient to restrict expression and render it rev-dependent rTHY1env:immunoglobulin fusion proteins were generated, in which only about one third of the total gene had the envelope codon usage. Expression levels of this construct were on an intermediate level, indicating that the rTHY-1env negative sequence element is not dominant over the immunoglobulin part. This fusion protein was not or only slightly rev-responsive, indicating that only genes almost completely suppressed can be rev-responsive.

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Another characteristic feature that was found in the codon frequency tables is a striking under representation of CpG triplets. In a comparative study of codon usage in E. coli, yeast, drosophila and primates it was shown that in a high number of analyzed primate genes the 8 least used codons contain all codons with the CpG dinucleotide sequence. Avoidance of codons containing this dinucleotide motif was also found in the sequence of other retroviruses. It seems plausible that the reason for under representation of CpG-bearing triplets has something to do with avoidance of gene silencing by methylation of CpG

cytosines. The expected number of CpG dinucleotides for HIV as a whole is about one fifth that expected on the basis of the base composition. This might indicate that the possibility of high expression is restored, and that the gene in fact has to be highly expressed at some point during viral pathogenesis.

The results presented herein clearly indicate that codon preference has a severe effect on protein levels, and suggest that translational elongation is controlling mammalian gene expression. However, other factors may play a role. First, abundance of not maximally loaded mRNA's in eukaryotic cells indicates that initiation is rate limiting for translation in at least some cases, since otherwise all transcripts would be completely covered by ribosomes. Furthermore, if ribosome stalling and subsequent mRNA degradation were the mechanism, suppression by rare codons could most likely not be reversed by any regulatory mechanism like the one presented herein. One possible explanation for the influence of both initiation and elongation on translational activity is that the rate of initiation, or access to ribosomes, is controlled in part by cues distributed throughout the RNA, such that the lentiviral codons predispose the RNA to accumulate in a pool of poorly initiated RNAs. However, this limitation need not be kinetic; for example, the choice of codons could influence the probability that a given translation product, once initiated, is properly completed. Under this mechanism, abundance of less favored codons would incur a significant cumulative probability of failure to complete the nascent polypeptide chain. The sequestered RNA would then be lent an improved rate of initiation by the action of rev. Since adenine residues are abundant in rev-responsive transcripts, it could be that RNA adenine methylation mediates this translational suppression.

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Detailed Procedures

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The following procedures were used in the above-described experiments.

Sequence Analysis

Sequence analyses employed the software developed by the University of Wisconsin Computer Group.

Plasmid constructions

Plasmid constructions employed the following methods. Vectors and insert DNA was digested at a concentration of 0.5 μ g/10 μ l in the appropriate restriction buffer for 1 - 4 hours (total reaction volume approximately 30 μ l). Digested vector was treated with 10% (v/v) of 1 μ g/ml calf intestine alkaline phosphatase for 30 min prior to gel electrophoresis. Both vector and insert digests (5 to 10 μ l each) were run on a 1.5% low melting agarose gel with TAE buffer. Gel slices containing bands of interest were transferred into a 1.5 ml reaction tube, melted at 65°C and directly added to the ligation without removal of the agarose. Ligations were typically done in a total volume of 25 μ l in 1x Low Buffer 1x Ligation Additions with 200-400 U of ligase, 1 μ l of vector, and 4 μ l of insert. When necessary, 5' overhanging ends were filled by adding 1/10 volume of 250 μ M dNTPs and 2-5 U of Klenow polymerase to heat inactivated or phenol extracted digests and incubating for approximately 20 min at room temperature. When necessary, 3' overhanging ends were filled by adding 1/10 volume of 2.5 mM dNTPs and 5-10 U of T4 DNA polymerase to heat inactivated or phenol extracted digests, followed by incubation at 37°C for 30 min. The following buffers were used in these reactions: 10x Low buffer (60 mM Tris HCl, pH 7.5, 60 mM MgCl₂, 50 mM NaCl, 4 mg/ml BSA, 70 mM β-mercaptoethanol, 0.02% NaN₃); 10x Medium buffer (60 mM Tris HCl, pH 7.5, 60 mM MgCl₂, 50 mM NaCl, 4 mg/ml BSA, 70 mM β - 5

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mercaptoethanol, 0.02% NaN₃); 10x High buffer (60 mM Tris HCl, pH 7.5, 60 mM MgCl₂, 50 mM NaCl, 4 mg/ml BSA, 70 mM β-mercaptoethanol, 0.02% NaN₃); 10x Ligation additions (1 mM ATP, 20 mM DTT, 1 mg/ml BSA, 10 mM spermidine); 50x TAE (2 M Tris acetate, 50 mM EDTA).

Oligonucleotide synthesis and purification

Oligonucleotides were produced on a Milligen 8750 synthesizer (Millipore). The columns were eluted with 1 ml of 30% ammonium hydroxide, and the eluted oligonucleotides were deblocked at 55°C for 6 to 12 hours. After deblockiong, 150 μ l of oligonucleotide were precipitated with 10x volume of unsaturated n-butanol in 1.5 ml reaction tubes, followed by centrifugation at 15,000 rpm in a microfuge. The pellet was washed with 70% ethanol and resuspended in 50 μ l of H₂0. The concentration was determined by measuring the optical density at 260 nm in a dilution of 1:333 (1 OD₂₆₀ = 30 μ g/ml).

The following oligonucleotides were used for construction of the synthetic gp120 gene (all sequences shown in this text are in 5' to 3' direction).

oligo 1 forward (Nhe1): cgc ggg cta gcc acc gag aag ctg (SEQ ID NO:1).

oligo 1: acc gag aag ctg tgg gtg acc gtg tac tac ggc gtg ccc gtg tgg aag ag ag gcc acc acc ctg ttc tgc gcc agc gac gcc aag gcg tac gac acc gag gtg cac aac gtg tgg gcc acc cag gcg tgc gtg ccc acc gac ccc aac ccc cag gag gtg gag ctc gtg aac gtg acc gag aac ttc aac at (SEQ ID NO:2).

oligo 1 reverse: cca cca tgt tgt tct tcc aca tgt tga agt tct c (SEQ ID NO:3).

oligo 2 forward: gac cga gaa ctt caa cat gtg gaa gaa caa cat (SEQ ID NO:4)

oligo 2 reverse (Pst1): gtt gaa gct gca gtt ctt cat ctc gcc gcc ctt (SEQ ID NO:6).

oligo 3 forward (Pst1): gaa gaa etg cag ett caa cat cac cac cag c (SEQ ID NO:7).

oligo 3: aac atc acc acc agc atc egc gac aag atg cag aag gag tac gec

ctg ctg tac aag etg gat atc gtg agc atc gac aac gac agc acc agc tac egc etg atc tec

tgc aac acc agc gtg atc acc eag gcc tgc ecc aag atc agc ttc gag ecc atc ecc atc

cac tac tgc gcc ecc gcc ggc ttc gcc (SEQ ID NO:8).

oligo 3 reverse: gaa ctt ctt gtc ggc ggc gaa gcc ggc ggg (SEQ ID NO:9).

oligo 4 forward: gcg ccc ccg ccg gct tcg cca tcc tga agt gca acg aca aga agt tc (SEQ ID NO:10)

oligo 4: gcc gac aag aag ttc agc ggc aag ggc agc tgc aag aac gtg agc acc gtg cag tgc acc cac ggc atc cgg ccg gtg gtg agc acc cag ctc ctg ctg aac ggc agc ctg gcc gag gag gtg gtg atc cgc agc gag aac ttc acc gac aac gcc aag acc atc atc gtg cac ctg aat gag agc gtg cag atc (SEQ ID NO:11)

oligo 4 reverse (Mlu1): agt tgg gac gcg tgc agt tga tct gca cgc tct c (SEQ ID NO:12).

oligo 5 forward (Mlu1): gag agc gtg cag atc aac tgc acg cgt ccc (SEQ ID NO:13).

oligo 5: aac tgc acg cgt ccc aac tac aac aag cgc aag cgc atc cac atc ggc ccc ggg cgc gcc ttc tac acc acc aag aac atc atc ggc acc atc ctc cag gcc cac tgc aac atc tct aga (SEQ ID NO:14).

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oligo 5 reverse: gtc gtt cca ctt ggc tct aga gat gtt gca (SEQ ID NO:15).

oligo 6 forward: gca aca tet eta gag eca agt gga acg ac (SEQ ID NO:16).

oligo 6: gcc aag tgg aac gac acc ctg cgc cag atc gtg agc aag ctg aag gag cag ttc aag aac aag acc atc gtg ttc ac cag agc agc ggc ggc gac ccc gag atc gtg atg cac agc ttc aac tgc ggc ggc (SEQ ID NO:17).

oligo 6 reverse (EcoR1): gca gta gaa gaa ttc gcc gcc gca gtt ga (SEQ ID NO:18).

oligo 7 forward (EcoR1):- tca act gcg gcg gcg aat tct tct act gc (SEQ ID NO:19).

oligo 7 reverse: gca gac cgg tga tgt tgc tgc tgc acc gga tct ggc cct c (SEQ ID NO:21).

oligo 8 forward: cga ggg cca gat ccg gtg cag cag caa cat cac cgg tct g (SEQ ID NO:22).

oligo 8: aac atc acc ggt ctg ctg ctg acc cgc gac ggc ggc aag gac acc gac acc aac gac acc gaa atc ttc cgc ccc ggc ggc ggc gac atg cgc gac aac tgg aga tct gag ctg tac aag tac aag gtg gtg acg atc gag ccc ctg ggc gtg gcc ccc acc aag gcc aag cgc cgc gtg gtg cag cgc gag aag cgc (SEQ ID NO:23).

oligo 8 reverse (Not1): cgc ggg cgg ccg ctt tag cgc ttc tcg cgc tgc acc ac (SEQ ID NO:24).

The following oligonucleotides were used for the construction of the ratTHY-1env gene.

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oligo 1 forward (BamH1/Hind3): cgc ggg gga tcc aag ctt acc atg att cca gta ata agt (SEQ ID NO:25).

oligo 1: atg aat cca gta ata agt ata aca tta tta tta agt gta tta caa atg agt aga gga caa aga gta ata agt tta aca gca tct tta gta aat caa aat ttg aga tta gat tgt aga cat gaa aat aat aca aat ttg cca ata caa cat gaa ttt tca tta acg (SEQ ID NO:26).

oligo 1 reverse (EcoR1/Mlu1): cgc ggg gaa ttc acg cgt taa tga aaa ttc atg ttg (SEQ ID NO:27).

oligo 2 forward (BamH1/Mlu1): cgc gga tcc acg cgt gaa aaa aaa aaa cat (SEQ ID NO:28).

oligo 2: cgt gaa aaa aaa aaa cat gta tta agt gga aca tta gga gta cca gaa cat aca tat aga agt aga gta aat ttg ttt agt gat aga ttc ata aaa gta tta aca tta gca aat ttt aca aca aaa gat gaa gga gat tat atg tgt gag (SEQ ID NO:29).

oligo 2 reverse (EcoR1/Sac1): cgc gaa ttc gag ctc aca cat ata atc tcc (SEQ ID NO:30).

oligo 3 forward (BamH1/Sac1): cgc gga tcc gag ctc aga gta agt gga caa (SEQ ID NO:31).

oligo 3 reverse (EcoR1/Not1): cgc gaa ttc gcg gcc gct tca taa act tat aaa atc (SEQ ID NO:33).

Polymerase Chain Reaction

Short, overlapping 15 to 25 mer oligonucleotides annealing at both ends were used to amplify the long oligonuclotides by polymerase chain reaction (PCR). Typical PCR conditions were: 35 cycles, 55°C annealing temperature, 0.2 sec extension time. PCR products were gel purified, phenol

extracted, and used in a subsequent PCR to generate longer fragments consisting of two adjacent small fragments. These longer fragments were cloned into a CDM7-derived plasmid containing a leader sequence of the CD5 surface molecule followed by a Nhe1/Pst1/Mlu1/EcoR1/BamH1 polylinker.

The following solutions were used in these reactions: 10x PCR buffer (500 mM KCl, 100 mM Tris HCl, pH 7.5, 8 mM MgCl₂, 2 mM each dNTP). The final buffer was complemented with 10% DMSO to increase fidelity of the Taq polymerase.

Small scale DNA preparation

Transformed bacteria were grown in 3 ml LB cultures for more than 6 hours or overnight. Approximately 1.5 ml of each culture was poured into 1.5 ml microfuge tubes, spun for 20 seconds to pellet cells and resuspended in 200 μ l of solution I. Subsequently 400 μ l of solution II and 300 μ l of solution III were added. The microfuge tubes were capped, mixed and spun for > 30 sec. Supernatants were transferred into fresh tubes and phenol extracted once. DNA was precipitated by filling the tubes with isopropanol, mixing, and spinning in a microfuge for > 2 min. The pellets were rinsed in 70 % ethanol and resuspended in 50 μ l dH20 containing 10 μ l of RNAse A. The following media and solutions were used in these procedures: LB medium (1.0 % NaCl, 0.5% yeast extract, 1.0% trypton); solution I (10 mM EDTA pH 8.0); solution II (0.2 M NaOH, 1.0% SDS); solution III (2.5 M KOAc, 2.5 M glacial aceatic acid); phenol (pH adjusted to 6.0, overlaid with TE); TE (10 mM Tris HCl, pH 7.5, 1 mM EDTA pH 8.0).

Large scale DNA preparation

One liter cultures of transformed bacteria were grown 24 to 36 hours (MC1061p3 transformed with pCDM derivatives) or 12 to 16 hours (MC1061 transformed with pUC derivatives) at 37°C in either M9 bacterial medium

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(pCDM derivatives) or LB (pUC derivatives). Bacteria were spun down in 1 liter bottles using a Beckman J6 centrifuge at 4,200 rpm for 20 min. The pellet was resuspended in 40 ml of solution I. Subsequently, 80 ml of solution II and 40 ml of solution III were added and the bottles were shaken semivigorously until lumps of 2 to 3 mm size developed. The bottle was spun at 4,200 rpm for 5 min and the supernatant was poured through cheesecloth into a 250 ml bottle.

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Isopropanol was added to the top and the bottle was spun at 4,200 rpm for 10 min. The pellet was resuspended in 4.1 ml of solution I and added to 4.5 g of cesium chloride, 0.3 ml of 10 mg/ml ethidium bromide, and 0.1 ml of 1% Triton X100 solution. The tubes were spun in a Beckman J2 high speed centrifuge at 10,000 rpm for 5 min. The supernatant was transferred into Beckman Quick Seal ultracentrifuge tubes, which were then sealed and spun in a Beckman ultracentrifuge using a NVT90 fixed angle rotor at 80,000 rpm for > 2.5 hours. The band was extracted by visible light using a 1 ml syringe and 20 gauge needle. An equal volume of dH₂O was added to the extracted material. DNA was extracted once with n-butanol saturated with 1 M sodium chloride, followed by addition of an equal volume of 10 M ammonium acetate/ 1 mM EDTA. The material was poured into a 13 ml snap tube which was tehn filled to the top with absolute ethanol, mixed, and spun in a Beckman J2 centrifuge at 10,000 rpm for 10 min. The pellet was rinsed with 70% ethanol and resuspended in 0.5 to 1 ml of H₂O. The DNA concentration was determined by measuring the optical density at 260 nm in a dilution of 1:200 (1 $OD_{260} = 50$ μ g/ml).

The following media and buffers were used in these procedures: M9 bacterial medium (10 g M9 salts, 10 g casamino acids (hydrolyzed), 10 ml M9 additions, 7.5 μ g/ml tetracycline (500 μ l of a 15 mg/ml stock solution), 12.5 μ g/ml ampicillin (125 μ l of a 10 mg/ml stock solution); M9 additions (10 mM

CaCl₂, 100 mM MgSO₄, 200 μg/ml thiamine, 70% glycerol); LB medium (1.0 % NaCl, 0.5 % yeast extract, 1.0 % trypton); Solution I (10 mM EDTA pH 8.0); Solution II (0.2 M NaOH 1.0 % SDS); Solution III (2.5 M KOAc 2.5 M HOAc)

Sequencing

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Synthetic genes were sequenced by the Sanger dideoxynucleotide method. In brief, 20 to 50 μ g double-stranded plasmid DNA were denatured in 0.5 M NaOH for 5 min. Subsequently the DNA was precipitated with 1/10 volume of sodium acetate (pH 5.2) and 2 volumes of ethanol and centrifuged for 5 min. The pellet-was-washed with 70% ethanol and resuspended at a concentration of 1 μ g/ μ l. The annealing reaction was carried out with 4 μ g of template DNA and 40 ng of primer in 1x annealing buffer in a final volume of 10 μ l. The reaction was heated to 65°C and slowly cooled to 37°C.

In a separate tube 1 μ l of 0.1 M DTT, 2 μ l of labeling mix, 0.75 μ l of dH₂0, 1 μ l of [35 S] dATP (10 μ Ci), and 0.25 μ l of SequenaseTM (12 U/ μ l) were added for each reaction. Five μ l of this mix were added to each annealed primer-template tube and incubated for 5 min at room temperature. For each labeling reaction 2.5 μ l of each of the 4 termination mixes were added on a Terasaki plate and prewarmed at 37°C. At the end of the incubation period 3.5 μ l of labeling reaction were added to each of the 4 termination mixes. After 5 min, 4 μ l of stop solution were added to each reaction and the Terasaki plate was incubated at 80°C for 10 min in an oven. The sequencing reactions were run on 5% denaturing polyacrylamide gel. An acrylamide solution was prepared by adding 200 ml of 10x TBE buffer and 957 ml of dH₂0 to 100 g of acrylamide:bisacrylamide (29:1). 5% polyacrylamide 46% urea and 1x TBE gel was prepared by combining 38 ml of acrylamide solution and 28 g urea. Polymerization was initiated by the addition of 400 μ l of 10% ammonium

peroxodisulfate and 60 μl of TEMED. Gels were poured using silanized glass plates and sharktooth combs and run in 1x TBE buffer at 60 to 100 W for 2 to 4 hours (depending on the region to be read). Gels were transferred to Whatman blotting paper, dried at 80 °C for about 1 hour, and exposed to x-ray film at room temperature. Typically exposure time was 12 hours. The following solutions were used in these procedures: 5x Annealing buffer (200 mM Tris HCl, pH 7.5, 100 mM MgCl₂, 250 mM NaCl); Labelling Mix (7.5 μM each dCTP, dGTP, and dTTP); Termination Mixes (80 μM each dNTP, 50 mM NaCl, 8 μM ddNTP (one each)); Stop solution (95% formamide, 20 mM EDTA, 0.05 % bromphenol blue, 0.05 % xylencyanol); 5x TBE (0.9 M Tris borate, 20 mM EDTA); Polyacrylamide solution (96.7 g polyacrylamide, 3.3 g bisacrylamide, 200 ml 1x TBE, 957 ml dH₂O).

RNA isolation

Cytoplasmic RNA was isolated from calcium phosphate transfected 293T cells 36 hours post transfection and from vaccinia infected Hela cells 16 hours post infection essentially as described by Gilman. (Gilman Preparation of cytoplasmic RNA from tissue culture cells. In Current Protocols in Molecular Biology, Ausubel et al., eds., Wiley & Sons, New York, 1992). Briefly, cells were lysed in 400 μ l lysis buffer, nuclei were spun out, and SDS and proteinase K were added to 0.2% and 0.2 mg/ml respectively. The cytoplasmic extracts were incubated at 37°C for 20 min, phenol/chloroform extracted twice, and precipitated. The RNA was dissolved in 100 μ l buffer I and incubated at 37°C for 20 min. The reaction was stopped by adding 25 μ l stop buffer and precipitated again.

The following solutions were used in this procedure: Lysis Buffer (TRUSTEE containing with 50 mM Tris pH 8.0, 100 mM NaCl, 5 mM MgCl₂, 0.5% NP40); Buffer I (TRUSTEE buffer with 10 mM MgCl₂, 1 mM DTT, 0.5

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 $U/\mu l$ placental RNAse inhibitor, 0.1 $U/\mu l$ RNAse free DNAse I); Stop buffer (50 mM EDTA 1.5 M NaOAc 1.0% SDS).

Slot blot analysis

For slot blot analysis 10 μ g of cytoplasmic RNA was dissolved in 50 μ l dH₂O to which 150 μ l of 10x SSC/18% formaldehyde were added. The solubilized RNA was then incubated at 65°C for 15 min and spotted onto with a slot blot apparatus. Radioactively labeled probes of 1.5 kb gp120IIIb and syngp120mn fragments were used for hybridization. Each of the two fragments was random labeled in a 50 μ l reaction with 10 μ l of 5x oligo-labeling buffer, 8 μ l of 2.5 mg/ml BSA, 4 μ l of [\propto ³²P]-dCTP (20 uCi/ μ l; 6000 Ci/mmol), and 5 U of Klenow fragment. After 1 to 3 hours incubation at 37°C 100 μ l of TRUSTEE were added and unincorporated [\approx^{32}P]-dCTP was eliminated using G50 spin column. Activity was measured in a Beckman beta-counter, and equal specific activities were used for hybridization. Membranes were prehybridized for 2 hours and hybridized for 12 to 24 hours at 42°C with 0.5 x 10⁶ cpm probe per ml hybridization fluid. The membrane was washed twice (5 min) with washing buffer I at room temperature, for one hour in washing buffer II at 65°C, and then exposed to x-ray film. Similar results were obtained using a 1.1 kb Not1/Sfi1 fragment of pCDM7 containing the 3 untranslated region. Control hybridizations were done in parallel with a random-labeled human beta-actin probe. RNA expression was quantitated by scanning the hybridized nitrocellulose membranes with a Magnetic Dynamics phosphorimager.

The following solutions were used in this procedure: 5x Oligo-labeling buffer (250 mM Tris HCl, pH 8.0, 25 mM MgCl₂, 5 mM β-mercaptoethanol, 2 mM dATP, 2 mM dGTP, mM dTTP, 1 M Hepes pH 6.6, 1 mg/ml hexanucleotides [dNTP]6); Hybridization Solution (.05 M sodium phosphate, 250 mM NaCl, 7% SDS, 1 mM EDTA, 5% dextrane sulfate, 50%

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formamide, 100 μ g/ml denatured salmon sperm DNA); Washing buffer I (2x SSC,

0.1% SDS); Washing buffer II (0.5x SSC, 0.1 % SDS); 20x SSC (3 M NaCl, 0.3 M Na₃citrate, pH adjusted to 7.0).

Vaccinia recombination

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Vaccinia recombination used a modification of the of the method described by Romeo and Seed (Romeo and Seed, Cell, 64: 1037, 1991). Briefly, CV1 cells at 70 to 90% confluency were infected with 1 to 3 \(\mu \)l of a wild-type vaccinia stock WR (2 x 108 pfu/ml) for 1 hour in culture medium without calf serum. After 24 hours, the cells were transfected by calcium phosphate with 25 μ g TKG plasmid DNA per dish. After an additional 24 to 48 hours the cells were scraped off the plate, spun down, and resuspended in a volume of 1 ml. After 3 freeze/thaw cycles trypsin was added to 0.05 mg/ml and lysates were incubated for 20 min. A dilution series of 10, 1 and 0.1 μ l of this lysate was used to infect small dishes (6 cm) of CV1 cells, that had been pretreated with 12.5 μ g/ml mycophenolic acid, 0.25 mg/ml xanthin and 1.36 mg/ml hypoxanthine for 6 hours. Infected cells were cultured for 2 to 3 days, and subsequently stained with the monoclonal antibody NEA9301 against gp120 and an alkaline phosphatase conjugated secondary antibody. Cells were incubated with 0.33 mg/ml NBT and 0.16 mg/ml BCIP in AP-buffer and finally overlaid with 1% agarose in PBS. Positive plaques were picked and resuspended in 100 μ l Tris pH 9.0. The plaque purification was repeated once. To produce high titer stocks the infection was slowly scaled up. Finally, one large plate of Hela cells was infected with half of the virus of the previous round. Infected cells were detached in 3 ml of PBS, lysed with a Dounce homogenizer and cleared from larger debris by centrifugation. VPE-8

recombinant vaccinia stocks were kindly provided by the AIDS repository,

Rockville, MD, and express HIV-1 IIIB gp120 under the 7.5 mixed early/late promoter (Earl et al., <u>I. Virol.</u>, 65:31, 1991). In all experiments with recombinant vaccina cells were infected at a multiplicity of infection of at least 10.

The following solution was used in this procedure:

AP buffer (100 mM Tris HCl, pH 9.5, 100 mM NaCl, 5 mM MgCl₂)

Cell culture

The monkey kidney carcinoma cell lines CV1 and Cos7, the human kidney carcinoma cell line 293T, and the human cervix carcinoma cell line. Hela were obtained from the American Tissue-Typing Collection and were maintained in supplemented IMDM. They were kept on 10 cm tissue culture plates and typically split 1:5 to 1:20 every 3 to 4 days. The following medium was used in this procedure:

Supplemented IMDM (90% Iscove's modified Dulbecco Medium, 10% calf serum, iron-complemented, heat inactivated 30 min 56°C, 0.3 mg/ml L-glutamine, 25 μ g/ml gentamycin 0.5 mM β -mercaptoethanol (pH adjusted with 5 M NaOH, 0.5 ml)).

Transfection

Calcium phosphate transfection of 293T cells was performed by
slowly adding and under vortexing 10 μg plasmid DNA in 250 μl 0.25 M
CaCl₂ to the same volume of 2x HEBS buffer while vortexing. After
incubation for 10 to 30 min at room temperature the DNA precipitate was
added to a small dish of 50 to 70% confluent cells. In cotransfection
experiments with rev, cells were transfected with 10 μg gp120IIIb,
gp120IIIbrre, syngp120mnrre or rTHY-1enveg1rre and 10 μg of pCMVrev or
CDM7 plasmid DNA.

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The following solutions were used in this procedure: 2x HEBS buffer (280 mM NaCl, 10 mM KCl, 1.5 mM sterile filtered); 0.25 mM CaCl₂ (autoclaved).

<u>Immunoprecipitation</u>

After 48 to 60 hours medium was exchanged and cells were incubated for additional 12 hours in Cys/Met-free medium containing 200 µCi of ³⁵S-translabel. Supernatants were harvested and spun for 15 min at 3000 rpm to remove debris. After addition of protease inhibitors leupeptin, aprotinin and PMSF to 2.5 µg/ml, 50 µg/ml, 100 µg/ml respectively, 1 ml of supernatant was incubated with either 10 µl of packed protein A sepharose alone (rTHY-1enveg1rre) or with protein A sepharose and 3 μg of a purified CD4/immunoglobulin fusion protein (kindly provided by Behring) (all gp120 constructs) at 4°C for 12 hours on a rotator. Subsequently the protein A beads were washed 5 times for 5 to 15 min each time. After the final wash 10 µl of loading buffer containing was added, samples were boiled for 3 min and applied on 7% (all gp120 constructs) or 10% (rTHY-lenveg1rre) SDS polyacrylamide gels (TRIS pH 8.8 buffer in the resolving, TRIS pH 6.8 buffer in the stacking gel, TRIS-glycin running buffer, Maniatis et al., supra 1989). Gels were fixed in 10% acetic acid and 10 % methanol, incubated with Amplify for 20 min, dried and exposed for 12 hours.

The following buffers and solutions were used in this procedure:

Wash buffer (100 mM Tris, pH 7.5, 150 mM NaCl, 5 mM CaCl₂, 1% NP-40);

5x Running Buffer (125 mM Tris, 1.25 M Glycin, 0.5% SDS); Loading buffer (10 % glycerol, 4% SDS, 4% β-mercaptoethanol, 0.02 % bromphenol blue).

Immunofluorescence

293T cells were transfected by calcium phosphate coprecipitation and analyzed for surface THY-1 expression after 3 days. After detachment

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with 1 mM EDTA/PBS, cells were stained with the monoclonal antibody OX-7 in a dilution of 1:250 at 4°C for 20 min, washed with PBS and subsequently incubated with a 1:500 dilution of a FITC-conjugated goat anti-mouse immunoglobulin antiserum. Cells were washed again, resuspended in 0.5 ml of a fixing solution, and analyzed on a EPICS XL cytofluorometer (Coulter).

The following solutions were used in this procedure:

PBS (137 mM NaCl, 2.7 mM KCl, 4.3 mM Na₂HPO₄, 1.4 mM KH₂PO₄, pH adjusted to 7.4); Fixing solution (2% formaldehyde in PBS).

ELISA

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—The concentration of gp120 in culture-supernatants was determined using CD4-coated ELISA plates and goat anti-gp120 antisera in the soluble phase. Supernatants of 293T cells transfected by calcium phosphate were harvested after 4 days, spun at 3000 rpm for 10 min to remove debris and incubated for 12 hours at 4°C on the plates. After 6 washes with PBS 100 μl of goat anti-gp120 antisera diluted 1:200 were added for 2 hours. The plates were washed again and incubated for 2 hours with a peroxidase-conjugated rabbit anti-goat IgG antiserum 1:1000. Subsequently the plates were washed and incubated for 30 min with 100 μl of substrate solution containing 2 mg/ml ophenylenediamine in sodium citrate buffer. The reaction was finally stopped with 100 μl of 4 M sulfuric acid. Plates were read at 490 nm with a Coulter microplate reader. Purified recombinant gp120IIIb was used as a control. The following buffers and solutions were used in this procedure: Wash buffer (0.1% NP40 in PBS); Substrate solution (2 mg/ml o-phenylenediamine in sodium citrate buffer).

EXAMPLE 2

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A Synthetic Green Fluorescent Protein Gene

The efficacy of codon replacement for gp120 suggests that replacing non-preferred codons with less preferred codons or preferred codons (and replacing less preferred codons with preferred codons) will increase expression in mammalian cells of other proteins, e.g., other eukaryotic proteins.

The green fluorescent protein (GFP) of the jellyfish Aequorea victoria (Ward, Photochem. Photobiol. 4:1, 1979; Prasher et al., Gene 111:229, 1992; Cody et al., Biochem. 32:1212, 1993) has attracted attention recently for its possible utility as a marker or reporter for transfection and lineage studies (Chalfie et al., Science 263:802, 1994).

Examination of a codon usage table constructed from the native coding sequence of GFP showed that the GFP codons favored either A or U in the third position. The bias in this case favors A less than does the bias of gp120, but is substantial. A synthetic gene was created in which the natural GFP sequence was re-engineered in much the same manner as for gp120 (FIG. 11; SEQ ID NO:40). In addition, the translation initiation sequence of GFP was replaced with sequences corresponding to the translational initiation consensus. The expression of the resulting protein was contrasted with that of the wild type sequence, similarly engineered to bear an optimized translational initiation consensus (FIG. 10B and FIG. 10C). In addition, the effect of inclusion of the mutation Ser 65-Thr, reported to improve excitation efficiency of GFP at 490 nm and hence preferred for fluorescence microscopy (Heim et al., Nature 373:663, 1995), was examined (FIG. 10D). Codon engineering conferred a significant increase in expression efficiency (an concomitant percentage of cells apparently positive for transfection), and the combination of

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the Ser 65-Thr mutation and codon optimization resulted in a DNA segment encoding a highly visible mammalian marker protein (FIG. 10D).

The above-described synthetic green fluorescent protein coding sequence was assembled in a similar manner as for gp120 from six fragments of approximately 120 bp each, using a strategy for assembly that relied on the ability of the restriction enzymes BsaI and BbsI to cleave outside of their recognition sequence. Long oligonucleotides were synthesized which contained portions of the coding sequence for GFP embedded in flanking sequences encoding EcoRI and BsaI at one end, and BamHI and BbsI at the other end. Thus, each oligonucleotide has the configuration EcoRI/BsaI/GFP fragment/BbsI/BamHI. The restriction site ends generated by the BsaI and BbsI sites were designed to yield compatible ends that could be used to join adjacent GFP fragments. Each of the compatible ends were designed to be unique and non-selfcomplementary. The crude synthetic DNA segments were amplified by PCR, inserted between EcoRI and BamHI in pUC9, and sequenced. Subsequently the intact coding sequence was assembled in a six fragment ligation, using insert fragments prepared with Bsal and BbsI. Two of six plasmids resulting from the ligation bore an insert of correct size, and one contained the desired full length sequence. Mutation of Ser65 to Thr was accomplished by standard PCR based mutagenesis, using a primer that overlapped a unique BssSI site in the synthetic GFP.

Codon optimization as a strategy for improved expression in mammalian cells

The data presented here suggest that coding sequence re-engineering may have general utility for the improvement of expression of mammalian and non-mammalian eukaryotic genes in mammalian cells. The results obtained here with three unrelated proteins: HIV gp120, the rat cell surface antigen Thy-

1 and green fluorescent protein from Aequorea victoria, and human Factor VIII (see below) suggest that codon optimization may prove to be a fruitful strategy for improving the expression in mammalian cells of a wide variety of eukaryotic genes.

EXAMPLE III

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BNSDOCID: <WO 9812207A1_IA>

Design of a Codon-Optimized Gene Expressing Human Factor VIII Lacking the Central B Domain

A synthetic gene was designed that encodes mature human Factor VIII lacking amino acid residues 760 to 1639, inclusive (residues 779 to 1658, inclusive, of the precursor). The synthetic gene was created by choosing codons corresponding to those favored by highly expressed human genes. Some deviation from strict adherence to the favored residue pattern was made to allow unique restriction enzyme cleavage sites to be introduced throughout the gene to facilitate future manipulations. For preparation of the synthetic gene the sequence was then divided into 28 segments of 150 basepairs, and a 29th segment of 161 basepairs.

The a synthetic gene expressing human Factor VIII lacking the central B domain was constructed as follows. Twenty-nine pairs of template oligonucleotides (see below) were synthesized. The 5' template oligos were 105 bases long and the 3' oligos were 104 bases long (except for the last 3' oligo, which was 125 residues long). The template oligos were designed so that each annealing pair composed of one 5' oligo and one 3' oligo, created a 19 basepair double-stranded regions.

To facilitate the PCR and subsequent manipulations, the 5' ends of the oligo pairs were designed to be invariant over the first 18 residues, allowing a common pair of PCR primers to be used for amplification, and allowing the same PCR conditions to be used for all pairs. The first 18 residues of each 5'

member of the template pair were cgc gaa ttc gga aga ccc (SEQ ID NO:110) and the first 18 residues of each 3' member of the template pair were: ggg gat cct cac gtc tca (SEQ ID NO:43).

Pairs of oligos were annealed and then extended and amplified by PCR in a reaction mixture as follows: templates were annealed at 200 μg/ml each in PCR buffer (10 mM Tris-HCl, 1.5 mM MgCl₂, 50 mM KCl, 100 μg/ml gelatin, pH 8.3). The PCR reactions contained 2 ng of the annealed template oligos, 0.5 μg of each of the two 18-mer primers (described below), 200 μM of each of the deoxynucleoside triphosphates, 10% by volume of DMSO and PCR buffer as supplied by Boehringer Mannheim Biochemicals, in a final volume of 50 μl. After the addition of Taq polymerase (2.5 units, 0.5 μl; Boehringer Mannheim Biochemicals) amplifications were conducted on a Perkin-Elmer Thermal Cycler for 25 cycles (94°C for 30 sec, 55°C for 30 sec, and 72°C for 30 sec). The final cycle was followed by a 10 minute extension at 72°C.

The amplified fragments were digested with EcoRI and BamHI (cleaving at the 5' and 3' ends of the fragments respectively) and ligated to a pUC9 derivative cut with EcoRI and BamHI.

Individual clones were sequenced and a collection of plasmids corresponding to the entire desired sequence was identified. The clones were then assembled by multifragment ligation taking advantage of restriction sites at the 3' ends of the PCR primers, immediately adjacent to the amplified sequence. The 5' PCR primer contained a BbsI site, and the 3' PCR primer contained a BsmBI site, positioned so that cleavage by the respective enzymes preceded the first nucleotide of the amplified portion and left a 4 base 5' overhang created by the first 4 bases of the amplified portion. Simultaneous digestion with BbsI and BsmBI thus liberated the amplified portion with unique 4 base 5' overhangs at each end which contained none of the primer sequences.

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In general these overhangs were not self-complementary, allowing multifragment ligation reactions to produce the desired product with high efficiency. The unique portion of the first 28 amplified oligonucleotide pairs was thereby 154 basepairs, and after digestion each gave rise to a 150 bp fragment with unique ends. The first and last fragments were not manipulated in this manner, however, since they had other restriction sites designed into them to facilitate insertion of the assembled sequence into an appropriate mammalian expression vector. The actual assembly process proceded as follows.

10 Assembly of the Synthetic Factor VIII Gene

Step 1: 29 Fragments Assembled to Form 10 Fragments.

The 29 pairs of oligonucleotides, which formed segments 1 to 29 when base-paired, are described below.

Plasmids carrying segments 1, 5, 9, 12, 16, 20, 24 and 27 were digested with EcoR1 and BsmBI and the 170 bp fragments were isolated; plasmids bearing segments 2, 3, 6, 7, 10, 13, 17, 18, 21, 25, and 28 were digested with BbsI and BsmBI and the 170 bp fragments were isolated; and plasmids bearing segments 4, 8, 11, 14, 19, 22, 26 and 29 were digested with EcoRI and BbsI and the 2440 bp vector fragment was isolated. Fragments bearing segments 1, 2, 3 and 4 were then ligated to generate segment "A"; fragments bearing segments 5, 6, 7 and 8 were ligated to generate segment "B"; fragments bearing segments 12, 13, and 14 were ligated to generate segment "C"; fragments bearing segments 16, 17, 18 and 19 were ligated to generate segment "D"; fragments bearing segments 20, 21 and 22 were ligated to generate segment "G"; fragments bearing segments 20, 21 and 25 were ligated to generate segment "G"; fragments bearing segments 24, 25 and 26 were ligated

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to generate segment "I"; and fragments bearing segments 27, 28 and 29 were ligated to generate segment "J".

Step 2: Assembly of the 10 resulting Fragments from Step 1 to Three Fragments.

Plasmids carrying the segments "A", "D" and "G" were digested with EcoRI and BsmBI, plasmids carrying the segments B, 15, 23, and I were digested with BbsI and BsmBI, and plasmids carrying the segments C, F, and J were digested with EcoRI and BbsI. Fragments bearing segments A, B, and C were ligated to generate segment "K"; fragments bearing segments D, 15, and F were ligated to generate segment "O"; and fragments bearing segments G, 23, I, and J were ligated to generate segment "P".

Step 3: Assembly of the Final Three Pieces.

The plasmid bearing segment K was digested with EcoRI and BsmBI, the plasmid bearing segment O was digested with BbsI and BsmBI, and the plasid bearing segment P was digested with EcoRI and BbsI. The three resulting fragments were ligated to generate segments.

Step 4: Insertion of the Synthetic Gene in a Mammalian Expression Vector.

The plasmid bearing segment S was digested with NheI and NotI and inserted between NheI and EagI sites of plasmid CD5lNEg1 to generate plasmid cd5lsf8b-.

Sequencing and Correction of the Synthetic Factor VIII Gene

After assembly of the synthetic gene it was discovered that there were two undesired residues encoded in the sequence. One was an Arg residue at 749, which is present in the GenBank sequence entry originating from Genentech but is not in the sequence reported by Genentech in the literature. The other was an Ala residue at 146, which should have been Pro. This

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mutation arose at an unidentified step subsequent to the sequencing of the 29 constituent fragments. The Pro749Arg mutation was corrected by incorporating the desired change in a PCR primer (ctg ctt ctg acg cgt gct ggg gtg gcg gga gtt; SEQ ID NO:44) that included the MluI site at position 2335 of the sequence below (sequence of HindIII to NotI segment) and amplifying between that primer and a primer (ctg ctg aaa gtc tcc agc tgc; SEQ ID NO:44) 5' to the SgrAI site at 2225. The SgrAI to MluI fragment was then inserted into the expression vector at the cognate sites in the vector, and the resulting correct sequence change verified by sequencing. The Pro146Ala mutation was corrected by incorporating the desired sequence change in an oligonucleotide (ggc agg tgc tta agg aga acg gcc cta tgg cca; SEQ ID NO:46) bearing the AfIII site at residue 504, and amplifying the fragment resulting from PCR reaction between that oligo and the primer having sequence cgt tgt tct tca tac gcg tct ggg gct cct cgg ggc (SEQ ID NO:109), cutting the resulting PCR fragment with AfIII and AvrII at (residue 989), inserting the corrected fragment into the expression vector and confirming the construction by sequencing. Construction of a Matched Native Gene Expressing Human Factor VIII Lacking the Central B Domain

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BNSDOCID: <WO 9812207A1 IA:

A matched Factor VIII B domain deletion expression plasmid having
the native codon sequence was constructed by introducing NheI at the 5' end of
the mature coding sequence using primer cgc caa ggg cta gcc gcc acc aga aga
tac tac ctg ggt (SEQ ID NO:47), amplifying between that primer and the primer
att cgt agt tgg ggt tcc tct gga cag (corresponding to residues 1067 to 1093 of the
sequence shown below), cutting with NheI and AfIII (residue 345 in the
sequence shown below) and inserting the resulting fragment into an
appropriately cleaved plasmid bearing native Factor VIII. The B domain
deletion was created by overlap PCR using ctg tat ttg atg aga acc g,

(corresponding to residues 1813 to 1831 below) and caa gac tgg tgg ggt ggc att aaa ttg ctt t (SEQ ID NO:48) (2342 to 2372 on complement below) for the 5' end of the overlap, and aat gcc acc cca cca gtc ttg aaa cgc ca (SEQ ID NO:49) (2352 to 2380 on sequence below) and cat ctg gat att gca ggg ag (SEQ ID NO:50) (3145 to 3164). The products of the two individual PCR reactions were then mixed and reamplified by use of the outermost primers, the resulting fragment cleaved by Asp718 (KpnI isoschizomer, 1837 on sequence below) and PflMI (3100 on sequence below), and inserted into the appropriately cleaved expression plasmid bearing native Factor VIII.

The complete sequence (SEQ-ID NO:41) of the native human factor VIII gene deleted for the central B region is presented in Figure 12. The complete sequence (SEQ ID NO:42) of the synthetic Factor VIII gene deleted for the central B region is presented in Figure 13.

Preparation and assay of expression plasmids

Two independent plasmid isolates of the native, and four independent isolates of the synthetic Factor VIII expression plasmid were separately propagated in bacteria and their DNA prepared by CsCl buoyant density centrifugation followed by phenol extraction. Analysis of the supernatants of COS cells transfected with the plasmids showed that the synthetic gene gave rise to approximately four times as much Factor VIII as did the native gene.

COS cells were then transfected with 5 µg of each factor VIII construct per 6 cm dish using the DEAE-dextran method. At 72 hours post-transfection, 4 ml of fresh medium containing 10% calf serum was added to each plated. A sample of media was taken from each plate 12 hr later. Samples were tested by ELISA using mouse anti-human factor VIII light chain monoclonal antibody and peroxidase-conjugated goat anti-human factor VIII

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polyclonal antibody. Purified human plasma factor VIII was used as a standard. Cells transfected with the synthetic Factor V111 gene construct expressed 138 ± 20.2 ng/ml (equivalent ng/ml non-deleted Factor VIII) of Factor VIII (n=4) while the cells transfected with the native Factor VIII gene expressed 33.5 ± 0.7 ng/ml (equivalent ng/ml non-deleted Factor VIII) of Factor VIII (n=2).

The following template oligonucleotides were used for construction of the synthetic Factor VIII gene.

---r1-bbs-1-for (gcta) -- --

cgc gaa ttc gga aga ccc gct agc cgc cac
ccg ccg cta cta cct ggg cgc cgt gga gct
gtc ctg gga cta cat gca gag cga cct ggg
cga gct ccc cgt gga (SEQ ID NO:51)

ggg gat cct cac gtc tca ggt ttt ctt gta 1 bam

15 cac cac gct ggt gtt gaa ggg gaa gct ctt

ggg cac gcg ggg ggg gaa gcg ggc gtc cac

ggg gag ctc gcc ca (SEQ ID NO:52)

rl bbs 2 for (aacc)

cgc gaa ttc gga aga ccc aac cct gtt cgt

20 gga gtt cac cga cca cct gtt caa cat tgc
caa gcc gcg ccc ccc ctg gat ggg cct gct
ggg ccc cac cat cca (SEQ ID NO:53)

2 r1

ggg gat cet cae gtc tea gtg cag get gae ggg gtg get ggc cat gtt ett cag ggt gat cae cae ggt gte gta cae ete gge etg gat ggt ggg gee cag ca (SEQ ID NO:54) 2 bam

5 r1 bbs 3 for (gcac)

cgc gaa ttc gga aga ccc gca cgc cgt ggg
cgt gag cta ctg gaa ggc cag cga ggg cgc
cga gta cga cga cca gac gtc cca gcg cga
gaa gga gga cga caa (SEQ ID NO:55)

3 r1

ggg gat cet cae gte tea get gge cat agg
gee gtt ete ett aag cae etg eea eae gta
ggt gtg get eee eee egg gaa eae ett gte
gte ete ett ete ge (SEQ ID NO:56)

3 bam

rl bbs 4 for (cagc)

cgc gaa ttc gga aga ccc cag cga ccc cct
gtg cct gac cta cag cta cct gag cca cgt
gga cct ggt gaa gga tct gaa cag cgg gct
gat cgg cgc cct gct (SEQ ID NO:57)

4 r1

ggg gat cet cae gte tea gaa eag eag gat

gaa ett gtg eag ggt etg ggt ttt ete ett

gge eag get gee ete geg aca eae eag eag

gge gee gat eag ee (SEQ ID NO:58)

4 bam

rl bbs 5 for (gttc)

cgc gaa ttc gga aga ccc gtt cgc cgt gtt
cga cga ggg gaa gag ctg gca cag cga gac
taa gaa cag cct gat gca gga ccg cga cgc
cgc cag cgc ccg cgc (SEQ ID NO:59)

ggg gat cet cae gte tea gtg gea gee gat
cag gee ggg cag get geg gtt cae gta gee
gtt aac ggt gtg cat ett ggg cea gge geg
-gge-get-gge-gge-gt (SEQ ID NO:60)

5 bam

rl bbs 6 for (ccac)

cgc gaa ttc gga aga ccc cca ccg caa gag
cgt gta ctg gca cgt cat cgg cat ggg cac
cac ccc tga ggt gca cag cat ctt cct gga
ggg cca cac ctt cct (SEQ ID NO:61)

6 rl

agt cag gat cet cac gte tea cag ggt etg gge
agt cag gaa ggt gat ggg get gat ete cag
get gge etg geg gtg gtt geg cac cag gaa
ggt gtg gee ete ca (SEQ ID NO:62)

6 bam

rl bbs 7 for (cctg)

cct agg cca gtt cct gct gtt ctg cca cat
cag cag cca cca gca cga cgg cat gga ggc
tta cgt gaa ggt gga (SEQ ID NO:63)

7 r1

ggg gat cet cae gte tea gte gte gte gta gte ete gge ete ete gtt gtt ett eat geg eag etg ggg ete ete ggg gea get gte eac ett eac gta age et (SEQ ID NO:64) 7 barr

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rl bbs 8 for (cgac)

cgc gaa ttc gga aga ccc cga cct gac cga cag cga gat gga tgt cgt acg ctt cga cga cga caa cag ccc cag ctt cat cca gat ccg cag cgt ggc caa-gaa (SEQ-ID NO:65)

8 rl

10

ggg gat cet cae gte tea tae tag egg gge gta gte cea gte ete ete ete gge gge gat gta gtg cae cea ggt ett agg gtg ett ett gge cae get geg ga (SEQ ID NO:66) 8 bam

rl bbs 9 for (agta)

cgc gaa ttc gga aga ccc agt act ggc ccc
cga cga ccg cag cta caa gag cca gta cct
gaa caa cgg ccc cca gcg cat cgg ccg caa
gta caa gaa ggt gcg (SEQ ID NO:67)

9 r1

ggg gat cet cae gte tea gag gat gee gga

cte gtg etg gat gge ete geg ggt ett gaa

agt ete gte ggt gta gge eat gaa geg eae

ett ett gta ett ge (SEQ ID NO:68)

9 bam

r1 bbs 10 for (cctc)

cgc gaa ttc gga aga ccc cct cgg ccc cct gct gta cgg cga ggt ggg cga cac cct gct gat cat ctt caa gaa cca ggc cag cag gcc cta caa cat cta ccc (SEQ ID NO:69) 10 r1

ggg gat cet cac gtc tca ctt cag gtg ctt
cac gcc ctt ggg cag gcg gcg gct gta cag
ggg gcg cac gtc ggt gat gcc gtg ggg gta
gat gtt gta ggg cc (SEQ ID NO:70)

10 bam

rl bbs 11 for (gaag)

cgc gaa ttc gga aga ccc gaa gga ctt ccc cat cct gcc cgg cga gat ctt caa gta caa gtg gac cgt gac cgt gga gga cgg ccc cac caa gag cga ccc ccg (SEQ ID NO:71)

11 r1

gga ggt cet cae gte tea gee gat eag tee
gga gge eag gte geg ete eat gtt eae gaa
get get gta gta geg ggt eag gea geg ggg
gte get ett ggt gg (SEQ ID NO:72)

11 bam

rl bbs 12 for (cggc)

cgc gaa ttc gga aga ccc cgg ccc cct gct
gat ctg cta caa gga gag cgt gga cca gcg
cgg caa cca gat cat gag cga caa gcg caa
cgt gat cct gtt cag (SEQ ID NO:73)

12 rl

ggg gat cet cae gte tea age ggg gtt ggg
cag gaa geg etg gat gtt ete ggt eag ata
cea get geg gtt ete gte gaa eae get gaa
cag gat eae gtt ge (SEQ ID NO:74)

12 bam

r1 bbs 13 for (cgct)

cgc gaa ttc gga aga ccc cgc tgg cgt gca gct gga aga tcc cga gtt cca ggc cag caa cat cat gca cag cat caa cgg cta cgt gtt cga-cag cct gca gct (SEQ-ID NO:75) 13 r1

ggg gat cet cae gte tea eag gaa gte ggt
ctg gge gee gat get eag gat gta eea gta
gge eac ete atg eag gea eac get eag etg
cag get gte gaa ea (SEQ ID NO:76)

13 bam

rl bbs 14 for (cctg)

cgc gaa ttc gga aga ccc cct gag cgt gtt
ctt ctc cgg gta tac ctt caa gca caa gat
ggt gta cga gga cac cct gac cct gtt ccc
ctt ctc cgg cga gac (SEQ ID NO:77)

14 r1

ggg gat cet cae gte tea gtt geg gaa gte

get gtt gtg gea gee eag aat eea eag gee
ggg gtt ete eat aga eat gaa eae agt ete
gee gga gaa ggg ga (SEQ ID NO:78)

14 bam

rl bbs 15 for (caac)

cgc gaa ttc gga aga ccc caa ccg cgg cat gac tgc cct gct gaa agt ctc cag ctg cga caa gaa cac cgg cga cta cta cga gga cag cta cga gga cat ctc (SEQ ID NO:79) 15 rl

ggg gat cet cae gte tea geg gtg geg gga gtt ttg gga gaa gga geg ggg ete gat gge gtt gtt ett gga eag eag gta gge gga gat gte ete gta get gt (SEQ ID NO:80) 15 bam

r1 bbs 16 for (ccgc)

cgc gaa ttc gga aga ccc ccg cag cac gcg tca gaa gca gtt caa cgc cac ccc ccc cgt gct gaa gcg cca cca gcg cga gat cac ccg cac cac cct gca aag (SEQ ID NO:81) 16 r1

ggg gat cet cae gte tea gat gte gaa gte
cte ett ett eat ete eae get gat ggt gte
gte gta gte gat ete ete etg gte get ttg
cag ggt ggt geg gg (SEQ ID NO:82)

16 bam

r1 bbs 17 for (catc)

20 cgc gaa ttc gga aga ccc cat cta cga cga
gga cga gaa cca gag ccc ccg ctc ctt cca
aaa gaa aac ccg cca cta ctt cat cgc cgc
cgt gga gcg cct gtg (SEQ ID NO:83)

17 r1

ggg gat cet cae gte tea etg ggg cae get gee get etg gge geg gtt geg eag gae gtg ggg get get get eat gee gta gte cea eag geg ete eae gge gg (SEQ ID NO:84) 17 bam

r1 bbs 18 for (ccag)

cgc gaa ttc gga aga ccc cca gtt caa gaa ggt ggt gtt cca gga gtt cac cga cgg cag ctt cac cca gcc cct gta ccg cgg cga gct gaa cga gca cct ggg (SEQ ID NO:85) 18 rl

ggg gat cct cac gtc tca ggc ttg gtt gcg
gaa ggt cac cat gat gtt gtc ctc cac ctc
ggc gcg gat gta ggg gcc gag cag gcc cag
gtg ctc gtt cag ct (SEQ ID NO:86)

18 bam

r1 bbs 19 for (agcc)

cgc gaa ttc gga aga ccc agc ctc ccg gcc
cta ctc ctt cta ctc ctc cct gat cag cta
cga gga gga cca gcg cca ggg cgc cga gcc
ccg caa gaa ctt cgt (SEQ ID NO:87)

19 r1

ggg gat cet cae gte tea ete gte ett ggt

20 ggg gge eat gtg gtg etg eae ett eea gaa
gta ggt ett agt ete gtt ggg ett eae gaa
gtt ett geg ggg et (SEQ ID NO:88)

19 bam

rl bbs 20 for (cgag)

cgc gaa ttc gga aga ccc cga gtt cga ctg
caa ggc ctg ggc cta ctt cag cga cgt gga
cct gga gaa gga cgt gca cag cgg cct gat
cgg ccc cct gct ggt (SEQ ID NO:89)

20 rl

ggg gat cet cae gte tea gaa cag gge aaa tte etg cae agt cae etg eet eee gtg ggg ggg gtt eag ggt gtt ggt gtg gea eae eag eag ggg gee gat ea (SEQ ID NO:90) 20 bam

10 r1 bbs 21 for (gttc)

cgc gaa ttc gga aga ccc gtt ctt cac cat ctt cga cga gac taa gag ctg gta ctt cac cga gaa cat gga gcg caa ctg ccg cgc ccc ctg caa cat cca gat (SEQ ID NO:91) 21 r1

ggg gat cet cac gtc tea cag ggt gtc cat
gat gta gcc gtt gat ggc gtg gaa gcg gta
gtt ctc ctt gaa ggt ggg atc ttc cat ctg
gat gtt gca ggg gg (SEQ ID NO:92)

21 bam

r1 bbs 22 for (cctg)

20 cgc gaa ttc gga aga ccc cct gcc cgg cct
ggt gat ggc cca gga cca gcg cat ccg ctg
gta cct gct gtc tat ggg cag caa cga gaa
cat cca cag cat cca (SEQ ID NO:93)

22 r1

ggg gat cet cae gte tea gta eag gtt gta
eag gge cat ett gta ete ete ett ett geg
eac ggt gaa aac gtg gee get gaa gtg gat
get gtg gat gtt et (SEQ ID NO:94)

22 bam

r1 bbs 23 for (gtac)

cgc gaa ttc gga aga ccc gta ccc cgg cgt
gtt cga gac tgt gga gat gct gcc cag caa
ggc cgg gat ctg gcg cgt gga gtg cct gat
cgg cga gca cct gca (SEQ ID NO:95)

23 rl

10 ggg gat cet cae gte tea get gge eat gee
cag ggg ggt etg gea ett gtt get gta eac
cag gaa eag ggt get eat gee gge gtg eag
gtg ete gee gat ea (SEQ ID NO:96)

23 bam

rl bbs 24 for (cagc)

cgc gaa ttc gga aga ccc cag cgg cca cat ccg cga ctt cca gat cac cgc cag cgg cca gta cgg cca gtg ggc tcc caa gct ggc ccg cct gca cta cag cgg (SEQ ID NO:97)

24 r1

ggg gat cet cae gte tea eat ggg gge eag

cag gte eac ett gat eea gga gaa ggg ete

ett ggt ega eea gge gtt gat get gee get

gta gtg eag geg gg (SEQ ID NO:98)

24 bam

rl bbs 25 for (catg)

cgc gaa ttc gga aga ccc cat gat cat cca cgg cat caa gac cca ggg cgc ccg cca gaa gtt cag cag cct gta cat cag cca gtt cat cat cat gta ctc tct (SEQ ID NO:99)

25 r1

ggg gat cct cac gtc tca gtt gcc gaa gaa cac cat cag ggt gcc ggt gct gtt gcc gcg gta ggt ctg cca ctt ctt gcc gtc tag aga gta cat gat gat ga (SEQ ID NO:100)

25 bam

rl bbs 26 for (caac)

cgc gaa ttc gga aga ccc caa cgt gga cag cag cgg cat caa gca caa cat ctt caa ccc ccc cat cat cgc ccg cta cat ccg cct gca ccc cac cca cta cag (SEQ ID NO:101) 26 r1

ggg gat cet cae gte tea gee eag ggg eat
get gea get gtt eag gte gea gee eat eag
ete eat geg eag ggt get geg gat get gta
gtg ggt ggg gtg ea (SEQ ID NO:102)

26 bam

r1 bbs 27 for (gggc)

caa ggc cat cag cga cgc cca gat cac cgc ctc cag cta ctt cac caa cat gtt cgc cac ctg gag ccc cag caa (SEQ ID NO:103)

27 rl

ggg gat cet cae gte tea cea etc ett ggg gtt gtt eac etg ggg geg eea gge gtt get geg gee etg eag gtg eag gge ett get ggg get eea ggt gg (SEQ ID NO:104) 27 bam

r1 bbs 28 for (gtgg)

cgc gaa ttc gga aga ccc gtg gct gca ggt gga ctt cca gaa aac cat gaa ggt gac tgg cgt gac cac cca ggg cgt caa gag cct gct gac cag cat gta cgt (SEQ ID NO:105) 28 r1

gaa gaa cag ggt ca ctg gtg gcc gtc ctg
gct gct gct gat cag gaa ctc ctt cac gta
cat gct ggt cag ca (SEQ ID NO:106)

28 bam

rl bbs 29 for (caag)

gtt cca ggg caa cca gga cag ctt cac acc ggt cgt gaa cag cct gga ccc ccc cct gct gac ccg cta cct gcg (SEQ ID NO:107) 29 rl

ggg gat cet cae gte tea geg gee get tea

gta cag gte etg gge ete gea gee cag cae

ete cat geg cag gge gat etg gtg cae cea

get etg ggg gtg gat geg cag gta geg ggt

cag ca (SEQ ID NO:108)

29 bam

The codon usage for the native and synthetic genes described above are presented in Tables 3 and 4, respectively.

TABLE 3: Codon Frequency of the Synthetic Factor VIII B Domain Deleted Gene

5	AA	Codon	Numb	er /100	00 Frac	ction
					,	
	Gly	GGG	7.00	4.82	0.09	
	Gly	GGA	1.00	0.69	0.01	
	Gly	GGT	0.00	0.00	0.00	
10	Gly	GGC	74.00	50.93	0.90	1,
	Glu	GAG	81.00	55.75	0.96	
	Glu	GAA	3.00	2.06	0.04	,1
	Asp	GAT	4.00	2.75	0.05	
15	Asp	GAC	78.00	53.68	0.95	
	Val	GTG	77.00	52.99	0.88	
	Val	GTA	2.00	1.38	0.02	
	Val	GTT	2.00	1.38	0.02	1
20	Val	GTC	7.00	4.82	0.08	
	Ala	GCG	0.00	0.00	0.00	
	Ala	GCA	0.00	0.00	0.00	
	Ala	GCT	3.00	2.06	0.04	
25	Ala	GCC	67.00	46.11	0.96	
	Arg	AGG	2.00_	1.38_	0.03	
	Arg	AGA	0.00	0.00	0.00	
	Ser	AGT	0.00	0.00	0.00	
30	Ser	AGC	97.00	66.76	0.81	
	Lys	AAG	75.00	51.62	0.94	
	Lys	AAA	5.00	3.44	0.06	
	Asn	AAT	0.00	0.00	0.00	
35	Asn	AAC	63.00	43.36	1.00	

	Met	ATG	43.00	29.59	1.00
	Ile	ATA	0.00	0.00	0.00
	Ile	ATT	2.00	1.38	0.03
5	Ile	ATC	72.00	49.55	0.97
	Thr	ACG	2.00		0.02
	Thr	ACA	1.00	0.69	0.01
	Thr	ACT	10.00	6.88	0.12
10	Thr	ACC	70.00	48.18	0.84
•	Trp	TGG	28.00	li li	1.00
	End	TGA	1.00	, 0.69	1.00
	Cys	TGT	1.00	0.69	0.05
15	Cys	TGC	18.00	12.39	0.95
	End	TAG	0.00	0.00	0.00
	End	TAA	0.00	0.00	0.00
	Tyr	TAT	2.00	1.38	0.03
20	Tyr	TAC	66.00	45.42	0.97
	Leu	TTG	0.00	0.00	0.00
	Leu	TTA	0.00	0.00	0.00
	Phe	TTT	1.00	0.69	0.01
25	Phe	TTC	76.00	52.31	0.99
	Ser	TCG	1.00	0.69	0.01
	Ser	TCA	0.00	0.00	0.00
	Ser	TCT	3.00	2.06	0.03
30	Ser	TCC	19.00	13.08	0.16
	Arg	CGG	1.00	0.69	0.01
	Arg	CGA	0.00	0.00	0.00
	Arg	CGT	1.00	0.69	0.01
35	Arg	CGC	69.00	47.49	0.95
	Gln	CAG	62.00	42.67	0.93
	Gln	CAA	5.00	3.44	0.07
	His	CAT	1.00	0.69	0.02
40	His	CAC	50.00	34.41	0.98

•	Leu	CTG	118.00	81.21	0.94
	Leu	CTA	3.00	2.06	0.02
	Leu	CTT	1.00	0.69	0.01
5	Leu	· CTC	3.00	2.06	0.02
•	Pro	CCG	4.00	2.75	0.05
	Pro	CCA	0.00	0.00	0.00
	Pro	CCT	3.00	2.06	0.04
10	Pro	CCC	68.00	46.80	0.91

TABLE⁻⁴: Codon Frequency Table of the Native Factor VIII B Domain Deleted Gene

15 '	AA	Codon	Numb	er /100	0 Fractio	n		
			· · · —					
	Gly	GGG	12.00	8.26	0.15			
	Gly	GGA	34.00	23.40	0.41			
	Gly	GGT	16.00	11.01	0.20		1	
20	Gly	GGC	20.00	13.76	0.24			
	Glu	GAG	33.00	22.71	0.39			
	Glu	GAA	51.00	35.10	0.61			
	Asp	GAT	55.00	37.85	0.67			
25	Asp	GAC	27.00	18.58	0.33	·	٠	
	Val	GTG	29.00	19.96	0.33			
	-Val	-GTA	-19.00	-13.08 -	0.22			
	Val	GTT	17.00	11.70	0.19			
30	Val	GTC	23.00	15.83	0.26			
	Ala	GCG	2.00	1.38	0.03			
	Ala	GCA	18.00	12.39	0.25			
	Ala	GCT	31.00	21.34	0.44			
35	Ala	GCC	20.00	13.76	0.28			

,					
1	Arg	AGG	18.00	12.39	0.25
	Arg	AGA	22.00	15.14	0.30
	Ser	AGT	22.00	15.14	0.18
	Ser	AGC	24.00	16.52	0.20
5					
	Lys	AAG	32.00	22.02	0.40
	Lys	AAA	48.00	33.04	0.60
	Asn	AAT	38.00	26.15	0.60
	Asn	AAC	25.00	17.21	0.40
10					
	Met	ATG	43.00	29.59	1.00
	Ile	ATA	13.00	8.95	0.18
	Ile	ATT	36.00	24.78	0.49
	Ile	ATC	25.00	17.21	0.34
15				*	
·	Thr	ACG	1.00	0.69	0.01
	Thr	ACA	23.00	15.83	0.28
, '	Thr	ACT	36.00	24.78	0.43
	Thr	ACC	23.00	15.83	0.28
20					
	Trp	TGG	28.00	19.27	1.00
	End	TGA	1.00	0.69	1.00
	Cys	TGT	7.00	4.82	0.37
	Cys	TGC	12.00	8.26	0.63
25					
	End	TAG	0.00	0.00	0.00
	End	TAA	0.00	0.00	0.00
	Tyr	TAT	41.00	28.22	0.60
	Tyr	TAC	27.00	18.58	0.40
30					
	Leu	TTG	20.00	13.76	0.16
	Leu	TTA	10.00	6.88	0.08
	Phe	TTT	45.00	30.97	0.58
	Phe	TTC	32.00	22.02	0.42
35					
	Ser	TCG	2.00	1.38	0.02
	Ser	TCA	27.00	18.58	0.22
	Ser	TCT	27.00	18.58	0.22
	Ser	TCC	18.00	12.39	0.15
40			•		

	Arg	CGG	6.00	4.13	0.08
	Arg	CGA	10.00	6.88	0.14
	Arg	CGT	7.00	4.82	0.10
	Arg	CGC	10.00	6.88	0.14
5	_	•		1	1
	Gln	CAG	42.00	28.91	0.63
	Gln	CAA	25.00	17.21	0.37
	His	CAT	28.00	19.27	0.55
	His	CAC	23.00	15.83	0.45
10			٠.	' 1	•
	Leu	CTG	36.00	24.78	0.29
•	Leu	CTA	15.00	10.32	0.12
	Leu	CTT	24.00	16.52	0.19
	Leu	CTC	20.00	13.76	0.16
15					
	Pro	CCG	1.00	0.69	0.01
	Pro	CCA	32.00	22.02	0.43
	Pro	CCT	26.00	17.89	0.35
	Pro	CCC	15.00	10.32	0.20
20		 			

<u>Use</u>

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30

The synthetic genes of the invention are useful for expressing the a protein normally expressed in mammalian cells in cell culture (e.g. for commercial production of human proteins such as hGH, TPA, Factor VIII, and Factor IX). The synthetic genes of the invention are also useful for gene therapy. For example, a synthetic gene encoding a selected protein can be introduced in to a cell which can express the protein to create a cell which can be administered to a patient in need of the protein. Such cell-based gene therapy techniques are well known to those skilled in the art, see, e.g., Anderson, et al., U.S. Patent No. 5,399,349; Mulligan and Wilson, U.S. Patent No. 5,460,959.

What is claimed is:

1. A synthetic gene encoding a protein normally expressed in an eukaryotic cell wherein at least one non-preferred or less preferred codon in a natural gene encoding said protein has been replaced by a preferred codon encoding the same amino acid, said synthetic gene being capable of expressing said protein at a level which is at least 110% of that expressed by said natural gene in an *in vitro* mammalian cell culture system under identical conditions.

- 2. The synthetic gene of claim 1 wherein said synthetic gene is capable of expressing said protein at a level which is at least 150% of that expressed by said natural gene in an *in vitro* cell culture system under identical conditions.
- 3. The synthetic gene of claim 1 wherein said synthetic gene is capable of expressing said protein at a level which is at least 200% of that expressed by said natural gene in an *in vitro* cell culture system under identical conditions.
- 4. The synthetic gene of claim 1 wherein said synthetic gene is capable of expressing said protein at a level which is at least 500% of that expressed by said natural gene in an *in vitro* cell culture system under identical conditions.
- 5. The synthetic gene of claim 1 wherein said synthetic gene comprises fewer than 5 occurrences of the sequence CG.
 - 6. The synthetic gene of claim 1 wherein at least 10% of the codons in said natural gene are non-preferred codons.

5

10

7. The synthetic gene of claim 1 wherein at least 50% of the codons in said natural gene are non-preferred codons.

- 8. The synthetic gene of claim 1 wherein at least 50% of the non-preferred codons and less preferred codons present in said natural gene have been replaced by preferred codons.
- 9. The synthetic gene of claim 1 wherein at least 90% of the non-preferred codons and less preferred codons present in said natural gene have been replaced by preferred codons.
- 10. The synthetic gene of claim 1 wherein said protein is normally expressed by a mammalian cell.
 - 11. The synthetic gene of claim 1 wherein said protein is a retroviral protein.
 - 12. The synthetic gene of claim 1 wherein said protein is a lentiviral protein.
- 13. The synthetic gene of claim 11 wherein said protein is an HIV protein.
 - 14. The synthetic gene of claim 13 wherein said protein is selected from the group consisting of gag, pol, and env.
 - 15. The synthetic gene of claim 13 wherein said protein is gp120.

- 16. The synthetic gene of claim 13 wherein said protein is gp160.
- 17. The synthetic gene of claim 1 wherein said protein is a human protein.
- 18. The synthetic gene of claim 1 wherein said human protein is
 5 Factor VIII.
 - 19. The synthetic gene of claim 1 wherein 20% of the codons are preferred codons.
 - 20. The synthetic gene of claim 18 wherein said gene has the coding sequence present in SEQ ID NO:42.
- 10 21. The synthetic gene of claim 1 wherein said protein is green fluorescent protein.
 - 22. The synthetic gene of claim 20 wherein said synthetic gene is capable of expressing said green fluorescent protein at a level which is at least 200% of that expressed by said natural gene in an *in vitro* mammalian cell culture system under identical conditions.
 - 23. The synthetic gene of claim 20 wherein said synthetic gene is capable of expressing said green fluorescent protein at a level which is at least 1000% of that expressed by said n atural gene in an *in vitro* mammalian cell culture system under identical conditions.

24. The synthetic gene of claim 21 having the sequence depicted in Figure 11 (SEQ ID NO:40).

- 25. An expression vector comprising the synthetic gene of claim 1.
- 5 26. The expression vector of claim 21, said expression vector being a mammalian expression vector.
 - 27. A mammalian cell harboring with the synthetic gene of claim 1.
- 28. A method for preparing a synthetic gene encoding a protein normally expressed by mammalian cells, comprising identifying non-preferred and less-preferred codons in the natural gene encoding said protein and replacing one or more of said non-preferred and less-preferred codons with a preferred codon encoding the same amino acid as the replaced codon.

Syngp120mn

	•				
1	CTCGAGATCC	ATTGTGCTCT	AAAGGAGATA	CCCGGCCAGA	CACCCTCACC
51	TGCGGTGCCC	AGCTGCCCAG	GCTGAGGCAA	GAGAAGGCCA	GAAACCATGC
101	CCATGGGGTC	TCTGCAACCG	CTGGCCACCT	TGTACCTGCT	GGGGATGCTG
151	GTCGCTTCCG	TGCTAGCCAC	CGAGAAGCTG	TGGGTGACCG	TGTACTACGG
201	CGTGCCCGTG	TGGAAGGAGG	CCACCACCAC	CCTGTTCTGC	GCCAGCGACG
251	CCAAGGCGTA	CGACACCGAG	GTGCACAACG	TGTGGGCCAC	CCAGGCGTGC
301	GTGCCCACCG	ACCCCAACCC	CCAGGAGGTG	GAGCTCGTGA	ACGTGACCGA
351	GAACTTCAAC	ATGTGGAAGA	ACAACATGGT	GGAGCAGATG	CATGAGGACA
401	TCATCAGCCT	GTGGGACCAG	AGCCTGAAGC	_CCTGCGTGAA	GCTGACCCCC
451	CTGTGCGTGA	CCCTGAACTG	CACCGACCTG	AGGAACACCA	CCAACACCAA
501	CAACAGCACC	GCCAACAACA	ACAGCAACAG	CGAGGGCACC	ATCAAGGGCG
551	GCGAGATGAA	GAACTGCAGC	TTCAACATCA	CCACCAGCAT	CCGCGACAAG
601	ATGCAGAAGG	AGTACGCCCT	GCTGTACAAG	CTGGATATCG	TGAGCATCGA
651	CAACGACAGC	ACCAGCTACC	GCCTGATCTC	CTGCAACACC	AGCGTGATCA
701	CCCAGGCCTG	CCCCAAGATC	AGCTTCGAGC	CCATCCCCAT	CCACTACTGC
751	GCCCCCGCCG	GCTTCGCCAT	CCTGAAGTGC	AACGACAAGA	AGTTCAGCGG
801	CAAGGGCAGC	TGCAAGAACG	TGAGCACCGT	GCAGTGCACC	CACGGCATCC
851	GGCCGGTGGT	GAGCACCCAG	CTCCTGCTGA	ACGGCAGCCT	GGCCGAGGAG
901	GAGGTGGTGA	TCCGCAGCGA	GAACTTCACC	GACAACGCCA	AGACCATCAT
951	CGTGCACCTG	AATGAGAGCG	TGCAGATCAA	CTGCACGCGT	CCCAACTACA
1001	ACAAGCGCAA	GCGCATCCAC	ATCGGCCCCG	GGCGCGCCTT	CTACACCACC
1051	AAGAACATCA	TCGGCACCAT	CCGCCAGGCC	CACTGCAACA	TCTCTAGAGC
1101	CAAGTGGAAC	GACACCCTGC	GCCAGATCGT	GAGCAAGCTG	AAGGAGCAGT
1151	TCAAGAACAA	GACCATCGTG	TTCAACCAGA	GCAGCGGCGG	CGACCCCGAG
1201	ATCGTGATGC	ACAGCTTCAA	CTGCGGCGGC	GAATTCTTCT	ACTGCAACAC
1251	CAGCCCCTG	TTCAACAGCA	CCTGGAACGG	CAACAACACC	TGGAACAACA
1301	CCACCGGCAG	CAACAACAAT	ATTACCCTCC	AGTGCAAGAT	CAAGCAGATC
1351	ATCAACATGT	GGCAGGAGGT	GGGCAAGGCC	ATGTACGCCC	CCCCCATCGA
1401	GGGCCAGATC	CGGTGCAGCA	GCAACATCAC	CGGTCTGCTG	CTGACCCGCG
1451	ACGGCGGCAA	GGACACCGAC	ACCAACGACA	CCGAAATCTT	CCGCCCCGGC

Fig. 1A

1501 GGCGGCGACA TGCGCGACAA CTGGAGATCT GAGCTGTACA AGTACAAGGT
1551 GGTGACGATC GAGCCCCTGG GCGTGGCCCC CACCAAGGCC AAGCGCCGCG

1601 TGGTGCAGCG CGAGAAGCGC TAAAGCGGCC GC

(SEQ ID NO: 34)

Fig. 1B

Syngp160mn

,					
1	ACCGAGAAGC	TGTGGGTGAC	CGTGTACTAC	GGCGTGCCCG	TGTGGAAGGA
51	GGCCACCACC	ACCCTGTTCT	GCGCCAGCGA	CGCCAAGGCG	TACGACACCG
101	AGGTGCACAA	CGTGTGGGCC	ACCCAGGCGT	GCGTGCCCAC	CGACCCCAAC
151	CCCCAGGAGG	TGGAGCTCGT	GAACGTGACC	GAGAACTTCA	ACATGTGGAA
201	GAACAACATG	GTGGAGCAGA	TGCATGAGGA	CATCATCAGC	CTGTGGGACC
251	AGAGCCTGAA	GCCCTGCGTG	AAGCTGACCC	CCCTGTGCGT	GACCCTGAAC
301	TGCACCGACC	TGAGGAACAC	CACCAACACC	AACAACAGCA	CCGCCAACAA
-351	-CAACAGCAAC	AGCGAGGGCA	CCATCAAGGG	CGGCGAGATG	AAGAACTGCA
401	GCTTCAACAT	CACCACCAGC	ATCCGCGACA	AGATGCAGAA	GGAGTACGCC
451	CTGCTGTACA	AGCTGGATAT	CGTGAGCATC	GACAACGACA	GCACCAGCTA
501	CCGCCTGATC	TCCTGCAACA	CCAGCGTGAT	CACCCAGGCC	TGCCCCAAGA
551	TCAGCTTCGA	GCCCATCCCC	ATCCACTACT	GCGCCCCCC	CGGCTTCGCC
601	ATCCTGAAGT	GCAACGACAA	GAAGTTCAGC	GGCAAGGGCA	GCTGCAAGAA
651	CGTGAGCACC	GTGCAGTGCA	CCCACGGCAT	CCGGCCGGTG	GTGAGCACCC
701	AGCTCCTGCT	GAACGGCAGC	CTGGCCGAGG	AGGAGGTGGT	GATCCGCAGC
751	GAGAACTTCA	CCGACAACGC	CAAGACCATC	ATCGTGCACC	TGAATGAGAG
801	CGTGCAGATC	AACTGCACGC	GTCCCAACTA	CAACAAGCGC	AAGCGCATCC
851	ACATCGGCCC	CGGCCGCCC	TTCTACACCA	CCAAGAACAT	CATCGGCACC
901	ATCCGCCAGG	CCCACTGCAA	CATCTCTAGA	GCCAAGTGGA	ACGACACCCT
951	GCGCCAGATC	GTGAGCAAGC	TGAAGGAGCA	GTTCAAGAAC	AAGACCATCG
1001	TGTTCAACCA	GAGCAGCGGC	GGCGACCCCG	AGATÇGTGAT	GCACAGCTTC
1051	AACTGCGGCG	GCGAATTCTT	CTACTGCAAC	ACCAGCCCCC	TGTTCAACAG
1101	CACCTGGAAC	GGCAACAACA	CCTGGAACAA	CACCACCGGC	AGCAACAACA
			ATCAAGCAGA		
1201	GTGGGCAAGG	CCATGTACGC	CCCCCCATC	GAGGGCCAGA	TCCGGTGCAG
1251	CAGCAACATC	ACCGGTCTGC	TGCTGACCCG	CGACGGCGGC	AAGGACACCG
1301	ACACCAACGA	CACCGAAATC	TTCCGCCCCG	GCGGCGGCGA	CATGCGCGAC
1351	AACTGGAGAT	CTGAGCTGTA	CAAGTACAAG	GTGGTGACGA	TCGAGCCCCT
1401	GGGCGTGGCC	CCCACCAAGG	CCAAGCGCCG	CGTGGTGCAG	CGCGAGAAGC
1451	GGGCCGCCAT	CGGCGCCCTG	TTCCTGGGCT	TCCTGGGGGC	GGCGGGCAGC

Fig. 1C

1501	ACCATGGGGG	CCGCCAGCGT	GACCCTGACC	GTGCAGGCCC	GCCTGCTCCT
1551	GAGCGGCATC	GTGCAGCAGC	AGAACAACCT	CCTCCGCGCC	ATCGAGGCCC
1601	AGCAGCATAT	GCTCCAGCTC	ACCGTGTGGG	GCATCAAGCA	GCTCCAGGCC
1651	CGCGTGCTGG	CCGTGGAGCG	CTACCTGAAG	GACCAGCAGC	TCCTGGGCTT
1701	CTGGGGCTGC	TCCGGCAAGC	TGATCTGCAC	CACCACGGTA	CCCTGGAACG
1751	CCTCCTGGAG	CAACAAGAGC	CTGGACGACA	TCTGGAACAA	CATGACCTGG
1801	ATGCAGTGGG	AGCGCGAGAT	CGATAACTAC	ACCAGCCTGA	TCTACAGCCT
1851	GCTGGAGAAG	AGCCAGACCC	AGCAGGAGAA	GAACGAGCAG	GAGCTGCTGG
1901	AGCTGGACAA	_GTGGGCGAGC	CTGTGGAACT	GGTTCGACAT	CACCAACTGG
1951	CTGTGGTACA	TCAAAATCTT	CATCATGATT	GTGGGCGCC	TGGTGGGCCT
2001	CCGCATCGTG	TTCGCCGTGC	TGAGCATCGT	GAACCGCGTG	CGCCAGGGCT
2051	ACAGCCCCCT	GAGCCTCCAG	ACCCGGCCCC	CCGTGCCGCG	CGGGCCCGAC
2101	CGCCCGAGG	GCATCGAGGA	GGAGGGCGGC	GAGCGCGACC	GCGACACCAG
2151	CGGCAGGCTC	GTGCACGGCT	TCCTGGCGAT	CATCTGGGTC	GACCTCCGCA
2201	GCCTGTTCCT	GTTCAGCTAC	CACCACCGCG	ACCTGCTGCT	GATCGCCGCC
2251	CGCATCGTGG	AACTCCTAGG	CCGCCGCGC	TGGGAGGTGC	TGAAGTACTG
2301	GTGGAACCTC	CTCCAGTATT	GGAGCCAGGA	GCTGAAGTCC	AGCGCCGTGA
2351	GCCTGCTGAA	CGCCACCGCC	ATCGCCGTGG	CCGAGGGCAC	CGACCGCGTG
2401	ATCGAGGTGC	TCCAGAGGGC	CGGGAGGGCG	ATCCTGCACA	TCCCCACCCG
2451	CATCCGCCAG	GGGCTCGAGA	GGGCGCTGCT	G (S	EQ ID NO: 35)

Fig. 1D

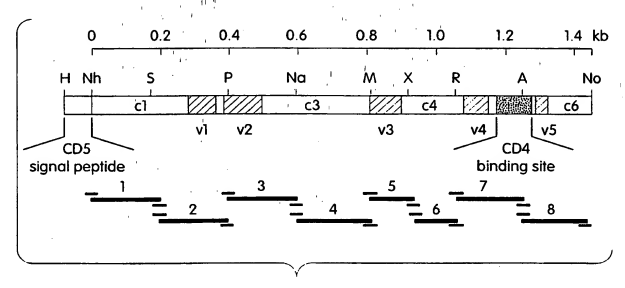


Fig. 2

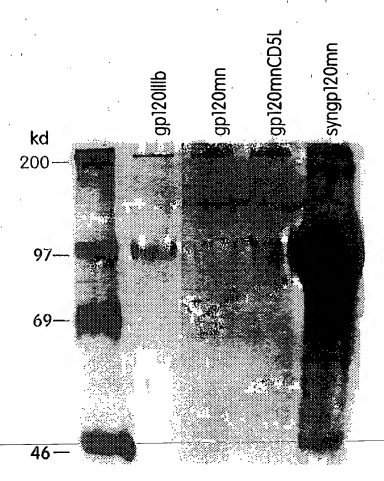


Fig. 3

SUBSTITUTE SHEET (RULE 26)

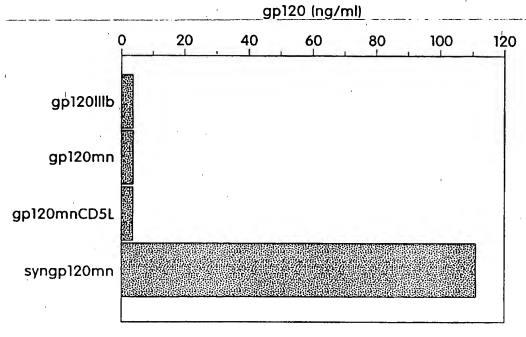


Fig. 4

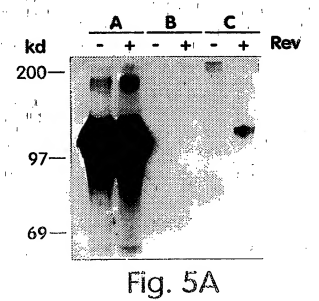
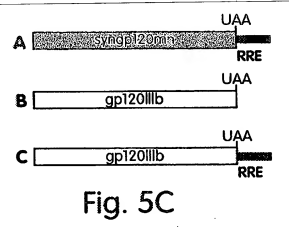
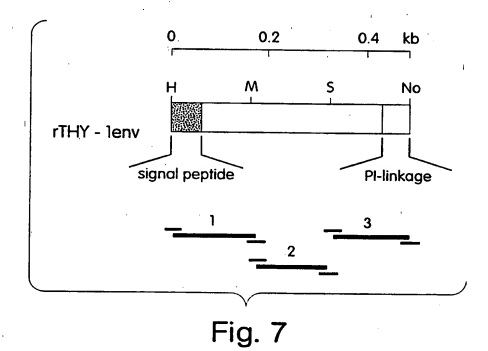




Fig. 5B



		' · · · · ·	. •	9/18				1
09	120	180	-240	300	360	420	480	T.
0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	G a t	22 22 X	ttg ctt L	999 990 6	ata atc I	aca act	agt tct s	
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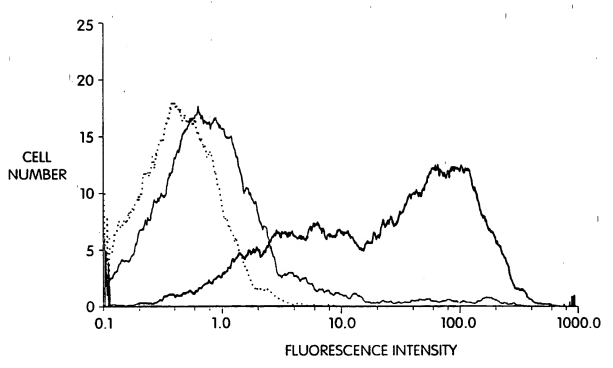


Fig. 8

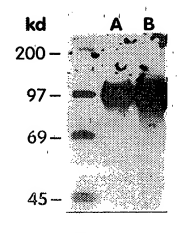
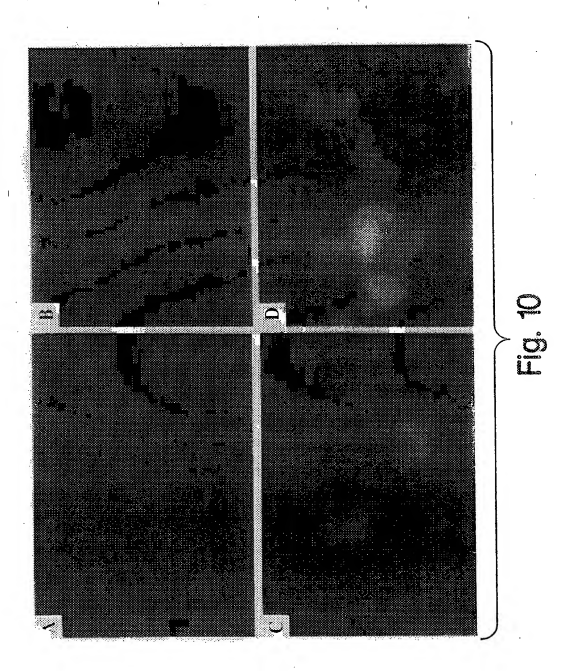


Fig. 9A

B Syngol 20 mm UAA

Fig. 9B

SUBSTITUTE SHEET (RULE 26).



SUBSTITUTE SHEET (RULE 26)

1	GAATTCACGC	GTAAGCTTGC	CGCCACCATG	GTGAGCAAGG	GCGAGGAGCT
51	GTTCACCGGG	GTGGTGCCCA	TCCTGGTCGA	GCTGGACGGC	GACGTGAACG
101	GCCACAAGTT	CAGCGTGTCC	GGCGAGGGCG	AGGGCGATGC	CACCTACGGC
151	AAGCTGACCC	TGAAGTTCAT	CTGCACCACC	GGCAAGCTGC	CCGTGCCCTG
201	GCCCACCCTC	GTGACCACCT	TCAGCTACGG	CGTGCAGTGC	TTCAGCCGCT
251	ACCCCGACCA	CATGAAGCAG	CACGACTTCT	TCAAGTCCGC	CATGCCCGAA
301	GGCTACGTCC	AGGAGCGCAC	CATCTTCTTC	AAGGACGACG	GCAACTACAA
351	GACCCGCGCC	GAGGTGAAGT	TCGAGGGCGA	CACCCTGGTG	AACCGCATCG
401	AGCTGAAGGG	CATCGACTTC	AAGGAGGACG	GCAACATCCT	GGGGCACAAG
451	CTGGAGTACA	ACTACAACAG	CCACAACGTC	TATATCATGG	CCGACAAGCA
501	GAAGAACGGC	ATCAAGGTGA	ACTTCAAGAT	CCGCCACAAC	ATCGAGGACG
551	GCAGCGTGCA	GCTCGCCGAC	CACTACCAGC	AGAACACCCC	CATCGGCGAC
601	GGCCCCGTGC	TGCTGCCCGA	CAACCACTAC	CTGAGCACCC	AGTCCGCCCT
651	GAGCAAAGAC	CCCAACGAGA	AGCGCGATCA	CATGGTCCTG	CTGGAGTTCG
701	TGACCGCCGC	CGGGATCACT	CACGGCATGG	ACGAGCTGTA	CAAGTAAAGC
751	GGCCGCGGAT	CC			

Fig. 11

	+			and the second s	
1	AAGCTTAAAC	CATGCCCATG	GGGTCTCTGC	AACCGCTGGC	CACCTTGTAC
51	CTGCTGGGGA	TGCTGGTCGC	TTCCGTGCTA	GCCGCCACCA	GAAGATACTA
101	CCTGGGTGCA	GTGGAACTGT	CATGGGACTA	TATGCAAAGT	GATCTCGGTG
151	AGCTGCCTGT	GGACGCAAGA	TTTCCTCCTA	GAGTGCCAAA	ATCTTTTCCA
201	TTCAACACCT	CAGTCGTGTA	CAAAAAGACT	CTGTTTGTAG	AATTCACGGA
251	TCACCTTTTC	AACATCGCTA	AGCCAAGGCC	ACCCTGGATG	GGTCTGCTAG
301	GTCCTACCAT.	CCAGGCTGAG	GTTTATGATA	CAGTGGTCAT	TACACTTAAG
351	AACATGGCTT	CCCATCCTGT	CAGTCTTCAT	GCTGTTGGTG	TATCCTACTG
401	GAAAGCTTCT	GAGGGAGCTG	AATATGATGA	TCAGACCAGT	CAAAGGGAGA
451	AAGAAGATGA	TAAAGTCTTC	CCTGGTGGAA	GCCATACATA	TGTCTGGCAG
501	GTCCTGAAAG	AGAATGGTCC	AATGGCCTCT	GACCCACTGT	GCCTTACCTA
551	CTCATATCTT	TCTCATGTGG	ACCTGGTAAA	AGACTTGAAT	TCAGGCCTCA
601	TTGGAGCCCT	ACTAGTATGT	AGAGAAGGGA [']	GTCTGGCCAA	GGAAAAGACA
651	CAGACCTTGC	ACAAATTTAT	ACTACTTTTT	GCTGTATTTG	ATGAAGGGAA
701	AAGTTGGCAC	TCAGAAACAA	AGAACTCCTT	GATGCAGGAT	AGGGATGCTG
751	CATCTGCTCG	GGCCTGGCCT	AAAATGCACA	CAGTCAATGG	TTATGTAAAC
801	AGGTCTCTGC	CAGGTCTGAT	TGGATGCCAC	AGGAAATCAG	TCTATTGGCA
851	TGTGATTGGA	ATGGGCACCA	CTCCTGAAGT	GCACTCAATA	TTCCTCGAAG
901	GTCACACATT	TCTTGTGAGG	AACCATCGCC	AGGCGTCCTT	GGAAATCTCG
951	CCAATAACTT	TCCTTACTGC	TCAAACACTC	TTGATGGACC	TTGGACAGTT
1001	TCTACTGTTT	TGTCATATCT	CTTCCCACCA	ACATGATGGC	ATGGAAGCTT
1051	ATGTCAAAGT	AGACAGCTGT	CCAGAGGAAC	CCCAACTACG	AATGAAAAT
1101	AATGAAGAAG	CGGAAGACTA	TGATGATGAT	CTTACTGATT	CTGAAATGGA
1151	TGTGGTCAGG	TTTGATGATG	ACAACTCTCC	TTCCTTTATC	CAAATTCGCT
1201	CAGTTGCCAA	GAAGCATCCT	AAAACTTGGG	TACATTACAT	TGCTGCTGAA
1251	GAGGAGGACT	GGGACTATGC	TCCCTTAGTC	CTCGCCCCCG	ATGACAGAAG
1301	TTATAAAAGT	CAATATTTGA	ACAATGGCCC	TCAGCGGATT	GGTAGGAAGT
1351	ACAAAAAAGT	CCGATTTATG	GCATACACAG	ATGAAACCTT	TAAGACTCGT
1401	GAAGCTATTC	AGCATGAATC	AGGAATCTTG	GGACCTTTAC	TTTATGGGGA
1451	AGTTGGAGAC	ACACTGTTGA	TTATATTTAA	GAATCAAGCA	AGCAGACCAT
1501	ATAACATCTA	CCCTCACGGA	ATCACTGATG	TCCGTCCTTT	GTATTCAAGG
1551	AGATTACCAA	AAGGTGTAAA	ACATTTGAAG	GATTTTCCAA	TTCTGCCAGG
1601	AGAAATATTC	AAATATAAAT	GGACAGTGAC	TGTAGAAGAT	GGGCCAACTA
1651	AATCAGATCC	TCGGTGCCTG	ACCCGCTATT	ACTCTAGTTT	CGTTAATATG
1701	GAGAGAGATC	TAGCTTCAGG	ACTCATTGGC	CCTCTCCTCA	TCTGCTACAA
1751	AGAATCTGTA	GATCAAAGAG	GAAACCAGAT	AATGTCAGAC	AAGAGGAATG
1801	TCATCCTGTT	TTCTGTATTT	GATGAGAACC	GAAGCTGGTA	CCTCACAGAG
1851	AATATACAAC	GCTTTCTCCC	CAATCCAGCT	GGAGTGCAGC	TTGAGGATCC
1901	AGAGTTCCAA	GCCTCCAACA	TCATGCACAG	CATCAATGGC	TATGTTTTTG
1951		GTTGTCAGTT	TGTTTGCATG	AGGTGGCATA	CTGGTACATT
2001		GAGCACAGAC	TGACTTCCTT		TCTCTGGATA
2051		CACAAAATGG	TCTATGAAGA		
2101			ATGTCGATGG		
2151			CTTTCGGAAC		
2201		AGTTGTGACA			
2251		TTCAGCATAC		AAAACAATGC	
2301			AAGACACCCT		
2351		CCACCAGTCT		TCAACGGGAA	
2401		GTCAGATCAA		ACTATGATGA	
2451		AGAAGGAAGA		TATGATGAGG	
2501		AGCTTTCAAA		ACACTATTTT	
2551		CTGGGATTAT		GCTCCCCACA	
2601		AGAGTGGCAG		TTCAAGAAAG	
2651	GGAATTTACT	GATGGCTCCT	TTACTCAGCC	CTTATACCGT	GGAGAACTAA

Fig. 12A

		• •			
2701	ATGAACATTT	GGGACTCCTG	GGGCCATATA	TAAGAGCAGA	AGTTGAAGAT
2751	AATATCATGG	TAACTTTCAG	AAATCAGGCC	TCTCGTCCCT	ATTCCTTCTA
2801	TTCTAGCCTT	ATTTCTTATG	AGGAAGATCA	GAGGCAAGGA	GCAGAACCTA
2851	GAAAAAACTT	TGTCAAGCCT	AATGAAACCA	AAACTTACTT	TTGGAAAGTG
2901	CAACATCATA	TGGCACCCAC	TAAAGATGAG	TTTGACTGCA	AAGCCTGGGC
2951	TTATTTCTCT	GATGTTGACC	TGGAAAAAGA	TGTGCACTCA	GGCCTGATTG
3001	GACCCCTTCT	GGTCTGCCAC	ACTAACACAC	TGAACCCTGC	TCATGGGAGA
3051	CAAGTGACAG	TACAGGAATT	TGCTCTGTTT	TTCACCATCT	TTGATGAGAC
3101	CAAAAGCTGG	TACTTCACTG	AAAATATGGA	AAGAAACTGC	AGGGCTCCCT
3151	GCAATATCCA	GATGGAAGAT	CCCACTTTTA	AAGAGAATTA	TCGCTTCCAT
3201	GCAATCAATG	GCTACATAAT	GGATACACTA	CCTGGCTTAG	TAATGGCTCA
3251	GGATCAAAGG	ATTCGATGGT	ATCTGCTCAG	CATGGGCAGC	AATGAAAACA
3301	TCCATTCTAT	TCATTTCAGT	GGACATGTGT	TCACTGTACG	AAAAAAGAG
3351	GAGTATAAAA	TGGCACTGTA	CAATCTCTAT	CCAGGTGTTT	TTGAGACAGT
3401	GGAAATGTTA	CCATCCAAAG	CTGGAATTTG	GCGGGTGGAA	TGCCTTATTG
3451	GCGAGCATCT	ACATGCTGGG	ATGAGCACAC	TTTTTCTGGT	GTACAGCAAT
3501	AAGTGTCAGA	CTCCCTGGG	AATGGCTTCT	GGACACATTA	GAGATTTTCA
3551	GATTACAGCT	TCAGGACAAT	ATGGACAGTG	GGCCCCAAAG	CTGGCCAGAC
3601	TTCATTATTC	CGGATCAATC	AATGCCTGGA	GCACCAAGGA	GCCCTTTTCT
3651	TGGATCAAGG	TGGATCTGTT	GGCACCAATG	ATTATTCACG	GCATCAAGAC
3701	CCAGGGTGCC	CGTCAGAAGT	TCTCCAGCCT	CTACATCTCT	CAGTTTATCA
3751	TCATGTATAG	TCTTGATGGG	AAGAAGTGGC	AGACTTATCG	AGGAAATTCC
3801	ACTGGAACCT	TAATGGTCTT	CTTTGGCAAT	GTGGATTCAT	CTGGGATAAA
3851	ACACAATATT	TTTAACCCTC	CAATTATTGC	TCGATACATC	CGTTTGCACC
3901	CAACTCATTA	TAGCATTCGC	AGCACTCTTC	GCATGGAGTT	GATGGGCTGT
3951	GATTTAAATA	GTTGCAGCAT	GCCATTGGGA	ATGGAGAGTA	AAGCAATATC
4001	AGATGCACAG	ATTACTGCTT	CATCCTACTT	TACCAATATG	TTTGCCACCT
4051	GGTCTCCTTC	AAAAGCTCGA	CTTCACCTCC	AAGGGAGGAG	TAATGCCTGG
4101	AGACCTCAGG	TGAATAATCC	AAAAGAGTGG	CTGCAAGTGG	ACTTCCAGAA
4151	GACAATGAAA	GTCACAGGAG	TAACTACTCA	GGGAGTAAAA	TCTCTGCTTA
4201	CCAGCATGTA	TGTGAAGGAG	TTCCTCATCT	CCAGCAGTCA	AGATGGCCAT
4251	CAGTGGACTC	TCTTTTTCA	GAATGGCAAA	GTAAAGGTTT	TTCAGGGAAA
4301	TCAAGACTCC	TTCACACCTG	TGGTGAACTC	TCTAGACCCA	CCGTTACTGA
4351	CTCGCTACCT	TCGAATTCAC	CCCCAGAGTT	GGGTGCACCA	GATTGCCCTG
4401	AGGATGGAGG	TTCTGGGCTG	CGAGGCACAG	GACCTCTACT	GAGGGTGGCC
4451	ACTGCAGCAC	CTGCCACTGC	CGTCACCTCT	CCCTCCTCAG	CTCCAGGGCA
4501	GTGTCCCTCC	CTGGCTTGCC	TTCTACCTTT	GTGCTAAATC	CTAGCAGACA
4551	CTGCCTTGAA	GCCTCCTGAA	TTAACTATCA	TCAGTCCTGC	ATTTCTTTGG
4601 4651	TGGGGGGCCA		TCCAATTTAA	CTTAACTCTT	ACCGTCGACC
4001	TCCACCCCA	ACCCCCCCCC			

Fig. 12B

		',			
1	AAGCTTAAAC	CATGCCCATG	GGGTCTCTGC	AACCGCTGGC	CACCTTGTAC
51	CTGCTGGGGA	TGCTGGTCGC	TTCCGTGCTA	GCCGCCACCC	GCCGCTACTA
101	CCTGGGCGCC	GTGGAGCTGT	CCTGGGACTA	CATGCAGAGC	GACCTGGGCG
151	AGCTCCCCGT	GGACGCCCGC	TTCCCCCCC	GCGTGCCCAA	GAGCTTCCCC
201	TTCAACACCA	GCGTGGTGTA	CAAGAAAACC	CTGTTCGTGG	AGTTCACCGA
251 '	CCACCTGTTC	AACATTGCCA	AGCCGCGCCC	CCCCTGGATG	GGCCTGCTGG
301	GCCCCACCAT	CCAGGCCGAG	GTGTACGACA	CCGTGGTGAT	CACCCTGAAG
351	AACATGGCCA	GCCACCCCGT	CAGCCTGCAC	GCCGTGGGCG	TGAGCTACTG
401	GAAGGCCAGC	GAGGGCGCCG	AGTACGACGA	CCAGACGTCC	CAGCGCGAGA
451	AGGAGGACGA	CAAGGTGTTC	CCGGGGGGGA	GCCACACCTA	CGTGTGGCAG
501	GTGCTTAAGG	AGAACGGCCC	TATGGCCAGC	GACCCCCTGT	GCCTGACCTA
551	CAGCTACCTG	AGCCACGTGG	ACCTGGTGAA	GGATCTGAAC	AGCGGGCTGA
601	TCGGCGCCCT	GCTGGTGTGT	CGCGAGGGCA	GCCTGGCCAA	GGAGAAAACC
651	CAGACCCTGC	ACAAGTTCAT	CCTGCTGTTC	GCCGTGTTCG	ACGAGGGGAA
701	GAGCTGGCAC	AGCGAGACTA	AGAACAGCCT	GATGCAGGAC	CGCGACGCCG
751	CCAGCGCCCG	CGCCTGGCCC	AAGATGCACA	CCGTTAACGG	CTACGTGAAC
801	CGCAGCCTGC	CCGGCCTGAT	CGGCTGCCAC	CGCAAGAGCG	TGTACTGGCA
851	CGTCATCGGC	ATGGGCACCA	CCCTGAGGT	GCACAGCATC	TTCCTGGAGG
901	GCCACACCTT	CCTGGTGCGC	AACCACCGCC	AGGCCAGCCT	GGAGATCAGC
951	CCCATCACCT	TCCTGACTGC	CCAGACCCTG	CTGATGGACC	TAGGCCAGTT
1001	CCTGCTGTTC	TGCCACATCA	GCAGCCACCA	GCACGACGGC	ATGGAGGCTT
1051	ACGTGAAGGT	GGACAGCTGC	CCCGAGGAGC	CCCAGCTGCG	CATGAAGAAC
1101	AACGAGGAGG	CCGAGGACTA	CGACGACGAC	CTGACCGACA	GCGAGATGGA
1151	TGTCGTACGC	TTCGACGACG	ACAACAGCCC	CAGCTTCATC	CAGATCCGCA
1201	GCGTGGCCAA	GAAGCACCCT	AAGACCTGGG	TGCACTACAT	CGCCGCCGAG
1251	GAGGAGGACT	GGGACTACGC	CCCGCTAGTA	CTGGCCCCCG	ACGACCGCAG
1301	CTACAAGAGC	CAGTACCTGA	ACAACGGCCC	CCAGCGCATC	GGCCGCAAGT
1351	ACAAGAAGGT	GCGCTTCATG	GCCTACACCG	ACGAGACTTT	CAAGACCCGC
1401	GAGGCCATCC	AGCACGAGTC	CGGCATCCTC	GGCCCCTGC	TGTACGGCGA
1451	GGTGGGCGAC	ACCCTGCTGA	TCATCTTCAA	GAACCAGGCC	AGCAGGCCCT
1501	ACAACATCTA	CCCCCACGGC	ATCACCGACG	TGCGCCCCCT	GTACAGCCGC
1551	CGCCTGCCCA	AGGGCGTGAA	GCACCTGAAG	GACTTCCCCA	TCCTGCCCGG
1601	CGAGATCTTC	AAGTACAAGT	GGACCGTGAC	CGTGGAGGAC	GGCCCCACCA
1651	AGAGCGACCC	CCGCTGCCTG	ACCCGCTACT	ACAGCAGCTT	CGTGAACATG
1701	GAGCGCGACC	TGGCCTCCGG	ACTGATCGGC	CCCCTGCTGA	TCTGCTACAA
1751	GGAGAGCGTG	GACCAGCGCG	GCAACCAGAT	CATGAGCGAC	AAGCGCAACG
1801	TGATCCTGTT	CAGCGTGTTC	GACGAGAACC	GCAGCTGGTA	TCTGACCGAG
1851	AACATCCAGC	GCTTCCTGCC	CAACCCCGCT	GGCGTGCAGC	TGGAAGATCC
1901	CGAGTTCCAG	GCCAGCAACA	TCATGCACAG	CATCAACGGC	TACGTGTTCG
1951	ACAGCCTGCA	GCTGAGCGTG	TGCCTGCATG	AGGTGGCCTA	CTGGTACATC
2001	CTGAGCATCG	GCGCCCAGAC	CGACTTCCTG	AGCGTGTTCT	TCTCCGGGTA
2051				CACCCTGACC	
2101	TCTCCGGCGA	GACTGTGTTC	ATGTCTATGG	AGAACCCCGG	CCTGTGGATT
2151	CTGGGCTGCC	ACAACAGCGA	CTTCCGCAAC	CGCGGCATGA	CTGCCCTGCT
2201				CGACTACTAC	
2251	ACGAGGACAT	CTCCGCCTAC	CTGCTGTCCA	AGAACAACGC	CATCGAGCCC
2301	CGCTCCTTCT	CCCAAAACTC	CCGCCACCCC	AGCACGCGTC	AGAAGCAGTT
2351		CCCCCGTGC		CCAGCGCGAG	
2401				ACTACGACGA	
2451				TACGACGAGG	
2501				CCACTACTTC	
2551				GCAGCCCCCA	-
2601				TTCAAGAAGG	
2651		GACGGCAGCT			
				-	

Fig. 13A

					1
2701	ACGAGCACCT	GGGCCTGCTC	GGCCCCTACA	TCCGCGCCGA	GGTGGAGGAC
2751	AACATCATGG	TGACCTTCCG	CAACCAAGCC	TCCCGGCCCT	ACTCCTTCTA
2801	CTCCTCCCTG	ATCAGCTACG	AGGAGGACCA	GCGCCAGGGC	GCCGAGCCCC
2851	GCAAGAACTT	CGTGAAGCCC	AACGAGACTA	AGACCTACTT	CTGGAAGGTG
2901	CAGCACCACA	TGGCCCCAC	CAAGGACGAG	TTCGACTGCA	AGGCCTGGGC
2951	CTACTTCAGC	GACGTGGACC	TGGAGAAGGA	CGTGCACAGC	GGCCTGATCG
3001	GCCCCTGCT	GGTGTGCCAC	ACCAACACCC	TGAACCCCCC	CCACGGGAGG
3051	CAGGTGACTG	TGCAGGAATT	TGCCCTGTTC	TTCACCATCT.	TCGACGAGAC
3101	TAAGAGCTGG	TACTTCACCG	AGAACATGGA	GCGCAACTGC	CGCGCCCCT
3151	GCAACATCCA	GATGGAAGAT	CCCACCTTCA	AGGAGAACTA	CCGCTTCCAC
3201	GCCATCAACG	GCTACATCAT	GGACACCCTG	CCCGGCCTGG	TGATGGCCCA
3251	GGACCAGCGC	ATCCGCTGGT	ACCTGCTGTC	TATGGGCAGC	AACGAGAACA
3301	TCCACAGCAT	CCACTTCAGC	GGCCACGTTT	TCACCGTGCG	CAAGAAGGAG
3351	GAGTACAAGA	TGGCCCTGTA	CAACCTGTAC	CCCGGCGTGT	TCGAGACTGT
3401	GGAGATGCTG	CCCAGCAAGG	CCGGGATCTG	GCGCGTGGAG	TGCCTGATCG
3451	GCGAGCACCT	GCACGCCGGC	ATGAGCACCC	TGTTCCTGGT	GTACAGCAAC
3501	AAGTGCCAGA	CCCCCTGGG	CATGGCCAGC	GGCCACATCC	GCGACTTCCA
3551	GATCACCGCC	AGCGGCCAGT	ACGGCCAGTG	GGCTCCCAAG	CTGGCCCGCC
3601	TGCACTACAG	CGGCAGCATC	AACGCCTGGT	CGACCAAGGA	GCCCTTCTCC
3651	TGGATCAAGG	TGGACCTGCT	GGCCCCCATG	ATCATCCACG	GCATCAAGAC
3701	CCAGGGCGCC	CGCCAGAAGT	TCAGCAGCCT	GTACATCAGC	CAGTTCATCA
3751	TCATGTACTC	TCTAGACGGC	AAGAAGTGGC	AGACCTACCG	CGGCAACAGC
3801	ACCGGCACCC	TGATGGTGTT	CTTCGGCAAC	GTGGACAGCA	GCGGCATCAA
3851	GCACAACATC	TTCAACCCCC	CCATCATCGC	CCGCTACATC	CGCCTGCACC
3901	CCACCCACTA	CAGCATCCGC	AGCACCCTGC	GCATGGAGCT	GATGGGCTGC
3951	GACCTGAACA	GCTGCAGCAT	GCCCCTGGGC	ATGGAGAGCA	AGGCCATCAG
4001	CGACGCCCAG	ATCACCGCCT	CCAGCTACTT	CACCAACATG	TTCGCCACCT
4051	GGAGCCCCAG	CAAGGCCCGC	CTGCACCTGC	AGGGCCGCAG	CAACGCCTGG
4101	CGCCCCAGG	TGAACAACCC	CAAGGAGTGG	CTGCAGGTGG	ACTTCCAGAA
4151	AACCATGAAG	GTGACTGGCG	TGACCACCCA	GGGCGTCAAG	AGCCTGCTGA
4201	CCAGCATGTA	CGTGAAGGAG	TTCCTGATCA	GCAGCAGCCA	GGACGGCCAC
4251	CAGTGGACCC	TGTTCTTCCA	AAACGGCAAG	GTGAAGGTGT	TCCAGGGCAA
4301	CCAGGACAGC	TTCACACCGG	TCGTGAACAG	CCTGGACCCC	CCCCTGCTGA
4351	CCCGCTACCT	GCGCATCCAC	CCCCAGAGCT	GGGTGCACCA	GATCGCCCTG
4401	CGCATGGAGG	TGCTGGGCTG	CGAGGCCCAG	GACCTGTACT	GAAGCGGCCG
4451	C				

Fig. 13B

International application No. PCT/US97/16639

IPC(6) US CL	SSIFICATION OF SUBJECT MATTER :C07H 21/04; C12P 21/02; C12N 15/11, 15/33, 15/45 :435/69.1, 70.1, 70.3, 172.3, 320.1; 536/23.1, 23.72, to International Patent Classification (IPC) or to both n	. 25.3
 	DS SEARCHED	actorial transfitation and in the
	ocumentation searched (classification system followed	by classification symbols)
	I I	
	435/69.1, 70.1, 70.3, 172.3, 320.1; 536/23.1, 23.72,	
Documental	ion searched other than minimum documentation to the e	xtent that such documents are included in the fields searched
		1.11
Electronic o	lata base consulted during the international search (nam	e of data base and, where practicable, search terms used)
BIOSIS, 1 search ter	EMBASE, MEDLINE, DERWENT ms: gene?, dna?, nucleic acid?, deoxyribonucleic?, syr	nthe?, prefer? non-prefer? codon?
c. Doc	UMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appr	ropriate, of the relevant passages Relevant to claim No.
X	WO 96/09378 A (THE GENERAL HO) 28 March 1996, abstract, page 1, line 20 lines 25-32, page 17, lines 27-39 and page 19, lines 27-39 and lines 27-39 and lines 27-39 and lines 29,	0-page 4, line 26, page 15,
A	SEETHARAM et al. Mistranslation Espression of the Protein in Escherichia Containing Low Frequency Codons. Comm. 30 August 1988. Vol. 155. N	coli Using a Synthetic Gene Biochem. Biophys. Res.
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Furti	ner documents are listed in the continuation of Box C.	See patent family annex.
•	ecial categories of cited documents: cument defining the general state of the art which is not considered	I'e later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
	be of particular relevance rlier document published on or after the international filing date	X° document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step
cil	ed to establish the publication date of another citation or other	when the document is taken alone Y* document of particular relevance; the claimed invention cannot be
*O" do	ecial reason (as specified) cument referring to an oral disclosure, use, exhibition or other cans	considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
"P" do		& document member of the same patent family
Date of the	actual completion of the international search	Date of mailing of the international search report
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